ONLINE REPORT

Improving Quality of Patient Data for Treatment of Multidrug- or Rifampin-Resistant Tuberculosis

Jonathon R. Campbell, Dennis Falzon, Fuad Mirzayev, Ernesto Jaramillo, Giovanni Battista Migliori, Carole D. Mitnick, Norbert Ndjeka, Dick Menzies

International policy for treatment of multidrug- and rifampin-resistant tuberculosis (MDR/RR TB) relies largely on individual patient data (IPD) from observational studies of patients treated under routine conditions. We prepared guidance on which data to collect and what measures could improve consistency and utility for future evidence-based recommendations. We highlight critical stages in data collection at which improvements to uniformity, accuracy, and completeness could add value to IPD quality. Through a repetitive development process, we suggest essential patient- and treatment-related characteristics that should be collected by prospective contributors of observational IPD in MDR/RR TB.

The treatment of multidrug- and rifampin-resistant tuberculosis (MDR/RR TB) is complex. Treatment requires a combination of multiple agents and often needs to be individualized, taking numerous considerations into account (1,2). Patients may have different concurrent conditions, such as HIV infection or diabetes; furthermore, the disease may vary in terms of extent, both in the lungs themselves (i.e., through presence of lung cavitation, bilateral disease, or both) and in other extrapulmonary sites (2). The pattern of additional resistances to other key agents used in second-line TB regimens may differ, depending on previous treatment received by the individual patient (either first-line or second-line medicines) and the epidemiologic setting (3,4). In different centers,

Author affiliations: McGill University, Montreal, Québec, Canada (J.R. Campbell, D. Menzies); Research Institute of the McGill University Health Centre, Montreal (J.R. Campbell, D. Menzies); World Health Organization, Geneva, Switzerland (D. Falzon, F. Mirzayev, E. Jaramillo); World Health Organization Collaborating Centre for Tuberculosis and Lung Diseases, Tradate, Italy (G.B. Migliori); Harvard Medical School, Boston, Massachusetts, USA (C.D. Mitnick); South African National Department of Health, Pretoria, South Africa (N. Ndjeka); Montréal Chest Institute, Montreal (D. Menzies)

DOI: https://doi.org/10.3201/eid2603.190997

the protocol for microbiologic monitoring may vary from none to monthly sputum smear microscopy and cultures with periodic drug-susceptibility testing during treatment, which, at times, continues after successful treatment to detect recurrence (2). The treatment given may be affected by the experience and expertise of the healthcare providers, as well as the cost of medicines and their availability (5). The use of adjunct therapies such as surgery (6), hospitalization, and patient support for treatment adherence (7), such as patient-centered directly observed therapy (DOT), also varies by program. The occurrence of adverse drug reactions to second-line TB drugs is common (1,8) and may be managed differently in different settings, particularly the permanent withdrawal of certain agents. All those factors result in wide variation in patient management and outcomes.

There is a shortage of high-quality randomized controlled trial (RCT) data for MDR/RR TB drugs (9), and currently available evidence is not adequately powered for patient outcomes (10–14). Although several notable RCTs evaluating standardized treatments are in the pipeline (15), no single regimen is likely to address the entire spectrum of clinical features that patients with MDR/RR TB have. This disease will largely require different treatment approaches individualized to the specific characteristics of the patient and the drug susceptibility profile of the strain.

Until the results of RCTs become available, new evidence for treatment of MDR/RR TB must be derived largely from observational studies. More than 150,000 MDR/RR TB patients initiate therapy each year worldwide, representing a wealth of potential data (16). These patients have an enormous diversity of clinical characteristics, many (e.g., pregnant women) are underrepresented in RCTs, and they are treated with widely varying regimens within health systems with different resources and capacities (17). This reflects the various scenarios in which global recommendations made by the World Health Orga-

nization (WHO) are expected to be applied and thus observational data can play a critical role in recommendation development.

Still, potential problems exist with use of observational data. The greatest are the potential for different forms of confounding and bias (18,19). This can be mitigated, at least partially, by careful adjustment for the many potential confounding factors, including age, prior treatment history, extent of drug resistance and disease, concurrent conditions, and treatment response (2). Adequate adjustment for confounders necessitates that information is accurately recorded for all patients treated, which is often not the case; missing data represents a second major potential limitation of observational data. Certain information may be missing for all patients in some centers, which could be the result of lack of capacity (e.g., radiography findings are missing because chest radiographs are not accessible) or the required information never being gathered or reported. Alternatively, other key data on determinants of patient outcomes, such as frequency and timing of regimen change, may be variably collected across studies. This may be caused by differences in the monitoring schedules, the data collection systems, and the medications used between studies and over time. At times, data collection may be directly related to determinants of outcome (e.g., length of QT-interval is more carefully measured and recorded in patients with multiple risks for cardiotoxicity) and can lead to measurement or ascertainment biases that are difficult to detect or mitigate appropriately.

Despite those problems, various studies have collected and pooled observational data, enabling individual patient meta-analyses (IPD-MAs). Since 2010, when WHO and other organizations started using GRADE for drug-resistant TB treatment guidelines (20), WHO recommendations on the type, composition, and duration of second-line TB regimens have been based largely on evidence from observational studies of patients treated under field conditions (21-25). Ahead of the WHO MDR/RR TB guideline update in 2018, a public call was made for contributors to report IPD conforming to certain criteria and a specific data dictionary (26). This call permitted including more recent programmatic data that may have never been published, increasing the breadth and relevance of the information available for study.

Overall, well-gathered, carefully documented, and complete observational datasets represent a valuable resource for assessing treatment regimens in MDR/RR TB. If efforts are made to safeguard the uniformity and quality of these data in terms of accuracy, consistency, and completeness, it is possible

to accrue sufficient information for large numbers patients treated for MDR/RR TB each year, and to generate evidence within 1–2 years to address critical questions, such as the optimal duration of the newly recommended all-oral MDR/RR TB regimen and the safety profile of new drugs (1). In response to our experiences with IPD management and analysis, most recently to update the WHO MDR/RR TB treatment recommendations in 2018 and 2019, and recognizing the urgent need for guidance, this article highlights how to improve the quality and completeness of future IPD for MDR/RR TB and provide guidance for researchers in other disease areas facing similar problems (27–30).

Aim and Scope of Guidance

Improving the completeness and quality of routinely collected data represents a relatively small marginal cost after all other expenditures incurred during care of patients with MDR/RR TB (31). Consolidating routinely reported data into high quality observational datasets and pooling these to perform multicentric IPD-MAs is a very attractive option to inform future MDR/RR TB treatment guidelines in the coming years, building on a proven track record (2,21–23,32–34).

The content of this guidance is meant for coordinators of MDR/RR TB treatment who intend to share their experience in patient care to the benefit of national and global treatment policy following several data-sharing principles (Table 1). This guidance is intended to instruct potential contributors on the utility of their potential observational IPD and aid them in subscribing to key quality and completeness measures to create a database with high quality IPD composed of key variables on patient demographics, clinical characteristics, treatment details and covariates, as well as treatment outcomes in MDR/RR TB patients, and contributing the IPD to a pooled data repository that can be shared internationally to allow for analysis that will inform future evidence-based treatment guidelines.

The guidance in this article was developed by 3 staff members of the WHO Global TB Programme (D.F., E.J., F.M.) involved in numerous iterations of the WHO MDR/RR TB guidelines and 5 methodologists, TB clinicians, and evidence reviewers (J.R.C., G.B.M., C.D.M., N.N., D.M.) involved in these and other guidelines. Four cycles of revisions took place, with successive discussions on key variables to collect, standardization of variable collection, and practical measures to suggest for completeness and quality. Although no one else was involved in writing the guidance, we acknowledge that we have benefited from the contribution and collective experience of

Table 1. Data-sharing principles for contributors of IPD for MDR/RR TB*

Principle	Additional notes
Data contributed to the IPD should be coded to remove identifying information.	 All names, the date of birth, address, telephone number, and other easily identifying personal information must be removed (e.g., national identification or health insurance numbers). Each participant contributed should be recoded with a new IPD identification number that is mapped to the original identification number retained by the contributing investigator, group, or program. Dates of events (e.g., treatment start, cultures, medication changes) should be retained in the sent participant data file. Other local rules for encoding and other data protection measures should be followed.
The contributing investigator, group, or program retains ownership of the data and should have permission to share them.	 The transfer of data for use in guideline development or other projects does not constitute transfer of ownership. Data contributors are free to withdraw their data at any time. Data must be contributed only if they are permitted by program or donor agencies. A data-sharing agreement will specify the details of the transfer of data; an example of a starting point for these data-sharing agreements is contained in the Appendix (https://wwwnc.cdc.gov/EID/article/26/3/19-0997-App1.pdf).
All transfers of data must clear ethics review.	 The institutional review board responsible for the bioethics of each contributed dataset should approve that the data can be shared. All anticipated uses of the data should be reviewed and approved by the institutional review board.
All uses of data are subject to oversight by the collaborative group.	 Ideally 1 individual is designated to liaise with the rest of the contributors of IPD to approve or deny use of their data for current or future analyses and be part of the oversight committee. The oversight committee reviews proposals for data use and sharing of data.
All data are held centrally in a secure data repository. *IPD, individual patient data; MDR, multidrug-resistant; MUHC, McGill Univer	 The IPD used for the development of MDR/RR TB treatment guidelines for the WHO and other entities has been held securely by the MUHC under Dick Menzies since 2010. The MUHC (now a WHO Collaborating Center) is expected to retain these responsibilities, pending approval of the oversight committee. Use of data held in this repository follows these principles, with bioethics approval and conforming to the current data sharing agreements signed.

many data contributors who provided data in the past and are acknowledged in publications of IPD-MA (2,21–23,26,32–34).

The Requirements for Observational Data

Several requirements exist to contributions of patient data. The first requirement is that the data are collected at centers that have the capacity to adequately gather the key information on all patients treated for MDR/RR TB. Centers should also have access to quality-assured medications in sufficient variety that they can treat patients with different drug susceptibility patterns. The centers should have adequate laboratory facilities to enable repeated microbiological testing throughout treatment, including initial and repeated drug susceptibility testing (DST) for all second-line TB medicines used at that center. Center staff should develop internal quality assurance protocols

and participate in external laboratory assessment programs to uphold the validity of their laboratory testing (35,36). These measures limit spurious conclusions being drawn about the influence of a medicine on outcomes resulting from exposure to ineffectual medication. The second requirement is that the program treats a relatively large number of patients with diverse demographic, clinical, and treatment characteristics. This policy avoids having patient series that are extreme outliers to the usual practice in a given setting. Nationwide representativeness is not to be expected, but reports of small patient series (e.g., <25) may be extreme outliers and may present challenges to pooling with other records for IPD-MA. However, we encourage reports of any size on subpopulations with limited available data, such as persons with extrapulmonary MDR/RR TB, pregnant women, children, and vulnerable populations. Finally, the

Table 2. Suggested steps to improve the accuracy and completeness of observational IPD	Table 2. Suggested ste	os to improve the accuracy	v and completeness of	observational IPD*
---	------------------------	----------------------------	-----------------------	--------------------

Suggested steps	Additional notes
Persons responsible for capture and entry of data into electronic databases should be appropriately trained.	 This includes obtaining a certificate in good clinical practice and training around the importance of confidentiality. This also includes training on the basics of MDR/RR TB, relevant national guidelines, what to collect, how to collect it, and the importance of accuracy in the capture of data. These principles can be reinforced with detailed guidance for data capture and the definitions of the variables collected at the point of capture (e.g., within the electronic system or within a document kept where data are captured).
Quality control measures (e.g., data safeguards) should be implemented to prevent implausible or "out-of-range" entries.	 A warning can be implemented for continuous variables falling outside plausible ranges (e.g., age outside 0–99 y). Drop-down lists can be created to reduce/remove need for free form data entry (e.g., including the most common extrapulmonary TB sites within the dropdown or limiting responses for HIV co-infection status to positive, negative, or not tested). Safeguards can be logical, which prevents certain data from being entered without a specific response in another section (e.g., CD4 and viral load cannot be filled in unless HIV co-infection status is positive).
Supervisors should have a standard quality assurance routine (e.g., perform routine follow-up for data accuracy of collected information).	 Supervisors should have simple algorithms developed to detect implausible information that defies inbuilt measures (e.g., patients reported to be receiving a medicine to which results from drug susceptibility testing show resistance). Complete checks should be run on at least 10% of records independently via dual extraction. These checks should be performed regularly and assessed by a supervisor with the goal of 95% accuracy. Corrective steps should be taken (e.g., further training, more comprehensive or routine checks of variables) when accuracy o data collection is an issue.
Concurrent checks for data completeness should be performed with assessments of accuracy.	 Reminders can be developed that automatically signal that certain variables are not completed each time a patient record is updated. In addition, preventing the "finalization" of a patient file until all variables are entered can be implemented—however, files should still be permitted to be saved, and other files opened and populated while patient files await finalization. Completeness of data is of utmost importance—high frequency of absence of certain information may necessitate exclusion of entire datasets from particular analyses for which these data are required.

center must adopt a quality-assured methodology for the study parameters and organization of data and respect ethics norms and standards for data collection, management, and use of data for research. This necessitates that the clinical data be entered in electronic format. Infrastructure must be in place to support electronic data collection, and personnel who are motivated and trained in data collection must be available. When possible, cross-checks should be performed between this electronic system and national vital statistics and laboratory registries, which provide information on long-term patient disposition.

Data Capture: Ensuring Accuracy and Completeness

Several practical measures should be undertaken to ensure that data are captured optimally. Upstream of the collection of data, efforts should be made to ensure the quality of these data, including quality assurance of diagnostic work and verification of patient demographic and clinical information with medical histories.

Transcription of data between systems (e.g., from a paper treatment card to an electronic database) is an eminent source of error. Many settings now have the capability to create an electronic medical record at the first encounter with the patient and access it again to prospectively update the details, either at subsequent patient visits or directly from the laboratory. The widespread availability of internet and desktop computers, laptops, tablets, or smartphones makes this feasible in many settings. This practice would have the advantages of improved completeness of patient files and avoidance of transcription and recall errors

when compared with other retrospective practices in data collection, such as periodic transfer of data from a paper treatment record during treatment, or after the treatment episode is completed. Within the electronic record system, anonymization procedures to limit the accidental disclosure of sensitive data are necessary. Various quality control measures can be built in to alert the user when implausible, inconsistent, or "outof-range"/nonstandardized values are entered, or if data are missing, prompting checks and corrections as necessary (Table 2). Finally, the database architecture of the health information system needs to allow for information from patient follow-up encounters to link up seamlessly to those of the initial record of the patient. A unique key in an electronic dataset limits the risk of duplicate records and avoids the need to re-enter identifiers of the patient and health center at each review. Many different packages have been successfully employed for this purpose, including opensource packages that bear no license fees for use and allow customization (37).

Description of Data Elements

This section highlights key items to capture within an electronic register (or database) for use in national or global analyses. The electronic medical record may contain other valuable information for programmatic management and policy making, such as health-related quality of life measurements, which may be of interest to programs, but which have not traditionally been used in analyses to date. The variables to be collected are those that are necessary to assess exposure (e.g., drugs, duration), potential confounders (e.g., concurrent conditions, resistance), response to treatment (e.g., microbiology, molecular biology, clinical signs and symptoms, and radiograph results), and adverse events (AEs). They also need to gather information that will be used to adjust observed effects by patient strata (e.g., by age, previous treatment history, or disease extent). A data dictionary defining variables and their preferred coding format is contained in the Appendix (http://wwwnc.cdc. gov/EID/article/26/3/19-0997-App1.pdf; the most up-to-date version of this data dictionary and accompanying tools and explanations are held at https:// www.mcgill.ca/tb/projects/mdr-tb-ipd-project). This list of variables is what is optimally preferred and what contributors should strive for; however, if certain data elements are missing from a patient series, the records may still be useful for specific analyses of safety or effectiveness. Further included in the Appendix are standard abbreviations for TB and antiretroviral drugs, standard system organ classes for

AEs (38), and standardized definitions for patient outcomes (39,40). We discuss variables that require further elaboration in the subsequent sections.

Initial (Baseline/Pretreatment)

Several baseline/pretreatment factors exist that affect the prognosis of patients with MDR/RR TB. Apart from typical demographic characteristics, complete collection of information on patients' habits and concurrent conditions is essential, as the true effect that many of these factors have on treatment outcomes is uncertain. Collection of CD4 counts, viral load, and antiretroviral therapy regimens in HIV-infected persons is essential; additional information on hepatitis B/C status, diabetes mellitus, and mental health disorders may also be useful. Although universally accepted definitions for smoking exist (41), this is not the case for alcohol consumption; contributors are encouraged to closely collect the alcohol-related variables in the data dictionary. The occurrence of cavitation and bilateral pulmonary disease is key to a better understanding of their effect on patient outcomes and to the classification of extent of disease. However, recording of radiologic findings in pulmonary MDR/ RR TB is not standardized between reporting centers and at times data are missing. For microbiological and DST results, several factors may compromise a program's ability to collect a sample exactly at treatment start. We suggest that baseline tests should be included only if they are performed on samples collected within 3 months before, or 1 month after, start of treatment. DST results should be reported for rifampin and for every medicine used in the regimen for which a WHO-approved laboratory method exists.

Treatment and Follow-Up Information

All measures that are repeated throughout treatment to inform treatment decisions and those that could affect treatment outcomes should be collected. It is perhaps most crucial to completely and accurately collect information regarding treatment type, duration, and composition. According to current standards, shorter MDR/RR TB regimens are those intended to last for ≤12 months, whereas longer regimens are intended to last for ≥18 months (1). Details for patients who had to transition from shorter to longer regimens must be reported. For each drug used in the regimen, ideally the day the drug was introduced into the regimen and the day the drug was permanently withdrawn (e.g., because of provider or patient decision or an adverse event) should be recorded. In programs in which this is not possible, new data elements can be added to the dictionary that would capture the patient's regimen

every 1-2 months, using standard abbreviations (Appendix). Adherence support, either in the form of inperson observation or with digital tools, is a common component of MDR/RR TB treatment. Data should be collected regarding its use and frequency. The data dictionary contains variables to record monthly follow-up sputum samples for smear microscopy and culture, with collection of culture results prioritized (1,42). Programs may also opt to simply report the date when each sputum sample was taken and the accompanying smear and culture result. Regardless of reporting choice, all results obtained should be recorded. Reporting of repeated DST is essential to detect acquired resistances; changes in the resistance patterns must be reported. Only thoracic surgery performed as an adjunctive therapy for MDR/RR TB should be reported.

The reporting of AEs in TB patients is highly valuable, but is often difficult to standardize. AEs of mild and moderate severity are very frequent in patients on TB treatment (1,8); including all of them in the IPD would be excessive. The AEs that should be entered and reported are drug-related AEs that are considered serious (43) or cases in which an agent is stopped for >48 hours by the provider because of a suspected or confirmed drug-related AE. In addition, information about whether the suspected or responsible agent is subsequently stopped permanently should be provided. Data in the "adverse event information" section should also be completed in the case of death that is suspected or confirmed to be drug-related. Characteristics of the AE that should be reported include the system or organ class affected, the agent(s) considered responsible, the severity, and the outcome. The severity should be graded using international standards, such as those of the National Cancer Institute (44) or other recommended scales (43,45). Centers may develop their own resources for the investigation and management of common AEs (e.g., by adapting the contents of manuals [46]).

Treatment Outcomes

End-of-treatment outcomes must be specified according to WHO standards to ensure uniformity. The set of definitions used must be specified, with preference currently given to 2013 criteria (1). Ideally, endpoint assignment would be systematically verified. Culture conversion (defined as the date of the first negative culture, when \geq 2 consecutive cultures, \geq 28 days apart, are negative) and culture reversion (defined as the date of the first positive culture, when \geq 2 consecutive cultures, \geq 28 days apart, are positive after culture conversion) should be reported (1). Recurrence

(because of true relapse or reinfection) information is valuable but scarce and difficult to collect because it requires follow-up after completion of treatment. The possibility to distinguish true relapses from a new infection among recurrences requires genotyping or sequencing that, to date, is done only in specialized laboratories, limiting its use in routine care (47). Monitoring patients for ≥12 months after successful completion of treatment would provide valuable information. If recurrence is monitored and reported, the exact duration of follow-up must be specified.

Discussion

We present a framework for observational data collection outlining key variables to collect to ensure uniformity in global MDR/RR TB patient data and provide practical measures to be taken to ensure data quality and completeness. National or regional TB programs, as well as operational research projects, patient series from a tertiary hospital, and other projects, could contribute their observational data through adoption of this guidance. However, wholesale adoption, especially by underresourced programs, will require support, in the form of funding and training, from donors, funding agencies, national programs, and others. The demonstrable value of IPD for developing WHO MDR/RR TB treatment guidelines (1,48,49) and continued need for quality IPD to tackle the MDR/RR TB epidemic underscore the importance of providing this support.

The strengths of this guidance are that it draws from our extensive experience in guideline revisions, IPD collection, and IPD-MA. Furthermore, our firsthand experience receiving retrospectively collected data conforming to the data dictionary (26) issued during the 2018 revision of the WHO MDR/RR TB guidelines provided valuable insight into barriers to data contribution. These barriers ranged from absence of crucial clinical and patient characteristics that were never recorded (and thus could not be retrospectively obtained) to difficulty in transcribing paper records of already-collected patient data into an electronic format. This guidance should provide motivation to programs to begin prospective data collection in a standardized electronic format, which is conducive to improvements in data completeness and quality. In addition, our experiences during guideline development highlighted key areas in which data were not routinely being collected (e.g., recurrence, acquired drug resistance) and populations for whom data were scarce. This guidance should encourage the collection of such data to help answer pressing questions in these domains and populations.

The primary limitation of this guidance is that it is an initial attempt to improve practices based on experience accumulated for a very particular subtype of patients with TB. The contents of the guidance will necessarily need to evolve to the ever-changing nature of MDR/RR TB treatment and the capacities of programs to adhere to it. Successive revisions will be informed as national TB programs and other end users begin to adopt this guidance and we gain experience receiving the outputs. Finally, certain variables, such as out-of-pocket costs, lost wages, specific toxicity-related measurements (e.g., electrocardiogram, brief peripheral neuropathy screens, audiometry, liver enzymes), emergence of mental health disorders, improvement or deterioration of quality of life, and emergence of AE that are not serious or do not result in medication termination, are not listed within our list of data elements. This information could be useful to patients, clinicians, and programs for specific studies, and thus could be added to local databases with care to avoid overloading data management.

Observational data will continue to play a critical role in the development of global MDR/RR TB treatment guidelines for the foreseeable future. Coordinating efforts to maximize the utility of provider experiences in MDR/RR TB is vital to improve the currently suboptimal outcomes of MDR/RR TB patients. This guidance is one key element toward achieving high-quality, comprehensive observational IPD moving forward.

Acknowledgments

We are grateful to J.J. Yim and Peter Cegielski, who provided a critical review of a draft of this online report.

This work was supported through a Unitaid grant to the World Health Organization (Unitaid-WHO Enabler grant 2018).

About the Author

Dr. Campbell is a postdoctoral fellow at McGill University, Montreal, Canada. His primary research interest is in tuberculosis and applying health economic, epidemiologic, and meta-analytical methods in its study.

References

- World Health Organization. WHO consolidated guidelines on drug-resistant tuberculosis treatment; 2019 [cited 2019 Nov 3]. https://www.who.int/tb/publications/2019/ consolidated-guidelines-drug-resistant-TB-treatment
- Collaborative Group for the Meta-Analysis of Individual Patient Data in MDR-TB treatment-2017, Ahmad N, Ahuja SD, Akkerman OW, Alffenaar J-WC, Anderson LF, et al. Treatment correlates of successful outcomes in pulmonary

- multidrug-resistant tuberculosis: an individual patient data meta-analysis. Lancet. 2018;392:821–34. https://doi.org/10.1016/S0140-6736(18)31644-1
- 3. Ruswa N, Mavhunga F, Roscoe JC, Beukes A, Shipiki E, van Gorkom J, et al. Second nationwide anti-tuberculosis drug resistance survey in Namibia. Int J Tuberc Lung Dis. 2019;23:858–64. https://doi.org/10.5588/ijtld.18.0526
- Centers for Disease Control and Prevention (CDC). Emergence of Mycobacterium tuberculosis with extensive resistance to second-line drugs – worldwide, 2000–2004. MMWR Morb Mortal Wkly Rep. 2006;55:301–5.
- Gotham D, Fortunak J, Pozniak A, Khoo S, Cooke G, Nytko FE III, et al. Estimated generic prices for novel treatments for drug-resistant tuberculosis. J Antimicrob Chemother. 2017;72:1243–52. https://doi.org/10.1093/jac/ dkw522
- Borisov SE, D'Ambrosio L, Centis R, Tiberi S, Dheda K, Alffenaar J-W, et al. Outcomes of patients with drugresistant-tuberculosis treated with bedaquiline-containing regimens and undergoing adjunctive surgery. J Infect. 2019;78:35–9. https://doi.org/10.1016/j.jinf.2018.08.003
- Law S, Daftary A, O'Donnell M, Padayatchi N, Calzavara L, Menzies D. Interventions to improve retention-in-care and treatment adherence among patients with drug-resistant tuberculosis: a systematic review. Eur Respir J. 2019;53:1801030. https://doi.org/10.1183/13993003.01030-2018
- Zhang Y, Wu S, Xia Y, Wang N, Zhou L, Wang J, et al. Adverse events associated with treatment of multidrugresistant tuberculosis in China: an ambispective cohort study. Med Sci Monit. 2017;23:2348–56. https://doi.org/10.12659/ MSM.904682
- Honeyborne I, Lipman M, Zumla A, McHugh TD. The changing treatment landscape for MDR/XDR-TB – can current clinical trials revolutionise and inform a brave new world? Int J Infect Dis. 2019;80S:S23–8. https://doi.org/ 10.1016/j.ijid.2019.02.006
- von Groote-Bidlingmaier F, Patientia R, Sanchez E, Balanag V Jr, Ticona E, Segura P, et al. Efficacy and safety of delamanid in combination with an optimised background regimen for treatment of multidrug-resistant tuberculosis: a multicentre, randomised, double-blind, placebo-controlled, parallel group phase 3 trial. Lancet Respir Med. 2019;7:249–59. https://doi.org/10.1016/ S2213-2600(18)30426-0
- Lee M, Lee J, Carroll MW, Choi H, Min S, Song T, et al. Linezolid for treatment of chronic extensively drugresistant tuberculosis. N Engl J Med. 2012;367:1508–18. https://doi.org/10.1056/NEJMoa1201964
- Diacon AH, Donald PR, Pym A, Grobusch M, Patientia RF, Mahanyele R, et al. Randomized pilot trial of eight weeks of bedaquiline (TMC207) treatment for multidrug-resistant tuberculosis: long-term outcome, tolerability, and effect on emergence of drug resistance. Antimicrob Agents Chemother. 2012;56:3271–6. https://doi.org/10.1128/ AAC.06126-11
- Tang S, Yao L, Hao X, Zhang X, Liu G, Liu X, et al. Efficacy, safety and tolerability of linezolid for the treatment of XDR-TB: a study in China. Eur Respir J. 2015;45:161–70. https://doi.org/10.1183/09031936.00035114
- Tang S, Yao L, Hao X, Liu Y, Zeng L, Liu G, et al. Clofazimine for the treatment of multidrug-resistant tuberculosis: prospective, multicenter, randomized controlled study in China. Clin Infect Dis. 2015;60:1361–7.
- Resist-TB. Clinical trial progress report; 2019 [cited 2019 May 15]. http://www.resisttb.org/?page_id=1602

ONLINE REPORT

- World Health Organization. Global tuberculosis report: 2018. 2019 [cited 2019 Nov 3]. https://apps.who.int/iris/bitstream/handle/10665/274453/9789241565646-eng.pdf
- World Health Organization. STOP TB. Contributing to health system strengthening: guiding principles for national tuberculosis programmes. 2008 [cited 2019 Nov 3]. https://www.who.int/tb/publications/tb-national-policy
- Franke MF, Rodriguez CA, Mitnick CD. Causal inference in tuberculosis treatment studies: bias considerations and data needs. Int J Tuberc Lung Dis. 2019;23:960–1. https://doi.org/ 10.5588/ijtld.19.0037
- Rodriguez CA, Mitnick CD, Franke MF. Value of observational data for multidrug-resistant tuberculosis. Lancet Infect Dis. 2019;19:930–1. https://doi.org/10.1016/ S1473-3099(19)30424-4
- Guyatt GH, Oxman AD, Vist GE, Kunz R, Falck-Ytter Y, Alonso-Coello P, et al.; GRADE Working Group. GRADE: an emerging consensus on rating quality of evidence and strength of recommendations. BMJ. 2008;336:924–6. https://doi.org/10.1136/bmj.39489.470347.AD
- Ahuja SD, Ashkin D, Avendano M, Banerjee R, Bauer M, Bayona JN, et al.; Collaborative Group for Meta-Analysis of Individual Patient Data in MDR-TB. Multidrug resistant pulmonary tuberculosis treatment regimens and patient outcomes: an individual patient data meta-analysis of 9,153 patients. PLoS Med. 2012;9:e1001300. https://doi.org/ 10.1371/journal.pmed.1001300
- Falzon D, Gandhi N, Migliori GB, Sotgiu G, Cox HS, Holtz TH, et al.; Collaborative Group for Meta-Analysis of Individual Patient Data in MDR-TB. Resistance to fluoroquinolones and second-line injectable drugs: impact on multidrug-resistant TB outcomes. Eur Respir J. 2013;42:156– 68. https://doi.org/10.1183/09031936.00134712
- Ahmad Khan F, Salim MAH, du Cros P, Casas EC, Khamraev A, Sikhondze W, et al. Effectiveness and safety of standardised shorter regimens for multidrug-resistant tuberculosis: individual patient data and aggregate data meta-analyses. Eur Respir J. 2017;50:1700061. https://doi.org/10.1183/13993003.00061-2017
- Fregonese F, Ahuja SD, Akkerman OW, Arakaki-Sanchez D, Ayakaka I, Baghaei P, et al. Comparison of different treatments for isoniazid-resistant tuberculosis: an individual patient data meta-analysis. Lancet Respir Med. 2018;6:265– 75. https://doi.org/10.1016/S2213-2600(18)30078-X
- Harausz EP, Garcia-Prats AJ, Law S, Schaaf HS, Kredo T, Seddon JA, et al.; Collaborative Group for Meta-Analysis of Paediatric Individual Patient Data in MDR-TB. Treatment and outcomes in children with multidrug-resistant tuberculosis: a systematic review and individual patient data meta-analysis. PLoS Med. 2018;15:e1002591. https://doi.org/ 10.1371/journal.pmed.1002591
- World Health Organization. Public call for individual patient data on treatment of rifampicin and multidrug-resistant (MDR/RR-TB) tuberculosis; 2018 [cited 2019 May 15]. http://www.who.int/tb/features_archive/public_call_ treatment_RR_MDR_TB
- Getahun H, Kittikraisak W, Heilig CM, Corbett EL, Ayles H, Cain KP, et al. Development of a standardized screening rule for tuberculosis in people living with HIV in resource-constrained settings: individual participant data meta-analysis of observational studies. PLoS Med. 2011;8:e1000391. https://doi.org/10.1371/ journal.pmed.1000391
- Schöttker B, Jorde R, Peasey A, Thorand B, Jansen EHJM, Groot L, et al.; Consortium on Health and Ageing: Network of Cohorts in Europe and the United States. Vitamin D and

- mortality: meta-analysis of individual participant data from a large consortium of cohort studies from Europe and the United States. BMJ. 2014;348(jun17 16):g3656. https://doi.org/10.1136/bmj.g3656
- 29. Ben-Shlomo Y, Spears M, Boustred C, May M, Anderson SG, Benjamin EJ, et al. Aortic pulse wave velocity improves cardiovascular event prediction: an individual participant meta-analysis of prospective observational data from 17,635 subjects. J Am Coll Cardiol. 2014;63:636–46. https://doi.org/10.1016/j.jacc.2013.09.063
- Morrison CS, Chen P-L, Kwok C, Baeten JM, Brown J, Crook AM, et al. Hormonal contraception and the risk of HIV acquisition: an individual participant data metaanalysis. PLoS Med. 2015;12:e1001778. https://doi.org/ 10.1371/journal.pmed.1001778
- 31. Yassin MA, Jaramillo E, Wandwalo E, Falzon D, Scardigli A, Kunii O, et al. Investing in a novel shorter treatment regimen for multidrug-resistant tuberculosis: to be repeated. Eur Respir J. 2017;49:1700081. https://doi.org/10.1183/13993003.00081-2017
- 32. Migliori GB, Sotgiu G, Gandhi NR, Falzon D, DeRiemer K, Centis R, et al.; Collaborative Group for Meta-Analysis of Individual Patient Data in MDR-TB. Drug resistance beyond extensively drug-resistant tuberculosis: individual patient data meta-analysis. Eur Respir J. 2013;42:169–79. https://doi.org/10.1183/09031936.00136312
- 33. Bastos ML, Hussain H, Weyer K, Garcia-Garcia L, Leimane V, Leung CC, et al.; Collaborative Group for Meta-analysis of Individual Patient Data in MDR-TB. Treatment outcomes of patients with multidrug-resistant and extensively drug-resistant tuberculosis according to drug susceptibility testing to first- and second-line drugs: an individual patient data meta-analysis. Clin Infect Dis. 2014;59:1364-74. https://doi.org/10.1093/cid/ciu619
- 34. Fox GJ, Mitnick CD, Benedetti A, Chan ED, Becerra M, Chiang C-Y, et al.; Collaborative Group for Meta-Analysis of Individual Patient Data in MDR-TB. Surgery as an adjunctive treatment for multidrug-resistant tuberculosis: an individual patient data metaanalysis. Clin Infect Dis. 2016;62:887–95. https://doi.org/10.1093/cid/ciw002
- 35. World Health Organization. WHO TB Supranational Reference Laboratory Network: TB diagnostics and laboratory strengthening; 2014 [cited 2019 Oct 16]. https://who.int/tb/areas-of-work/laboratory/srl-network
- 36. World Health Organization. Framework for conducting reviews of tuberculosis programmes. 2014 [cited 2019 Nov 3]. https://www.who.int/tb/publications/framework-tb-programme-reviews
- 37. District Health Information Software. DHIS2 overview; 2019 [cited 2019 May 15]. https://www.dhis2.org/overview
- International Council for Harmonisation of Technical Requirements for Pharmaceuticals for Human Use. MedDRA: introductory guide version 21.0. Geneva: The Council; 2018.
- Laserson KF, Thorpe LE, Leimane V, Weyer K, Mitnick CD, Riekstina V, et al. Speaking the same language: treatment outcome definitions for multidrug-resistant tuberculosis. Int J Tuberc Lung Dis. 2005;9:640–5.
- World Health Organization. Definitions and reporting framework for tuberculosis: 2013 revision. 2014 [cited 2019 Nov 3]. https://www.who.int/tb/publications/definitions
- Bernaards CM, Twisk JW, Snel J, Van Mechelen W, Kemper HC. Is calculating pack-years retrospectively a valid method to estimate life-time tobacco smoking? A comparison between prospectively calculated pack-years and retrospectively calculated pack-years. Addiction. 2001;

- 96:1653-61. https://doi.org/10.1046/j.1360-0443.2001. 9611165311.x
- Kurbatova EV, Cegielski JP, Lienhardt C, Akksilp R, Bayona J, Becerra MC, et al. Sputum culture conversion as a prognostic marker for end-of-treatment outcome in patients with multidrug-resistant tuberculosis: a secondary analysis of data from two observational cohort studies. Lancet Respir Med. 2015;3:201–9. https://doi.org/10.1016/ S2213-2600(15)00036-3
- Halleux CM, Falzon D, Merle C, Jaramillo E, Mirzayev F, Olliaro P, et al. The World Health Organization global aDSM database: generating evidence on the safety of new treatment regimens for drug-resistant tuberculosis. Eur Respir J. 2018;51:1701643. https://doi.org/10.1183/13993003.01643-2017
- 44. National Cancer Institute. Common Terminology Criteria for Adverse Events (CTCAE), v4.0; 2009 [cited 2019 Nov 3]. https://www.eortc.be/services/doc/ctc/ CTCAE_4.03_2010-06-14_QuickReference_5x7.pdf
- USAID, Systems for Improved Access to Pharmaceuticals and Services. Pharmacovigilance Monitoring System (PViMS). Arlington (TX): Management Sciences for Health; 2015.

- International Council of Nurses, Curry International Tuberculosis Center. Nursing guide for managing side effects to drug-resistant TB treatment. Geneva: The Council; 2018.
- 47. Migliori GB; Global Tuberculosis Network (GTN). Evolution of programmatic definitions used in tuberculosis prevention and care. Clin Infect Dis. 2019;68:1787–9. https://doi.org/10.1093/cid/ciy990
- 48. World Health Organization. WHO treatment guidelines for drug-resistant tuberculosis. 2016 update. 2016 [cited 2019 Nov 3]. https://apps.who.int/iris/bitstream/handle/10665/250125/9789241549639-eng.pdf
- World Health Organization. Guidelines for the programmatic management of drug-resistant tuberculosis – 2011 update.
 2011 [cited 2019 Nov 3]. https://www.who.int/tb/ challenges/mdr/programmatic_guidelines_for_mdrtb

Address for correspondence: Dick Menzies, McGill University, Office 3D.58, 5252 Boulevard de Maisonneuve O, Montréal, QB H4A 3S5, Canada; email: dick.menzies@mcgill.ca



EMERGING INFECTIOUS DISEASES

September 2016

Antimicrobial Resistance

- Co-Infections in Visceral Pentastomiasis, Democratic Republic of the Congo
- Multistate US Outbreak of Rapidly Growing Mycobacterial Infections Associated with Medical Tourism to the Dominican Republic, 2013–2014
- Virulence and Evolution of West Nile Virus, Australia, 1960–2012
- Phylogeographic Evidence for 2 Genetically Distinct Zoonotic Plasmodium knowlesi Parasites, Malaysia
- Hemolysis after Oral Artemisinin Combination Therapy for Uncomplicated *Plasmodium falciparum* Malaria
- Enterovirus D68 Infection in Children with Acute Flaccid Myelitis, Colorado, USA, 2014
- Middle East Respiratory Syndrome Coronavirus Transmission in Extended Family, Saudi Arabia, 2014
- Exposure-Specific and Age-Specific Attack Rates for Ebola Virus Disease in Ebola-Affected Households, Sierra Leone
- Outbreak of *Achromobacter xylosoxidans* and *Ochrobactrum anthropi* Infections after Prostate Biopsies, France, 2014
- Probable Rabies Virus Transmission through Organ Transplantation, China, 2015

- Human Babesiosis, Bolivia, 2013
- Assessment of Community Event–Based Surveillance for Ebola Virus Disease, Sierra Leone, 2015
- Cutaneous Melioidosis Cluster Caused by Contaminated Wound Irrigation Fluid
- Possible Role of Fish and Frogs as Paratenic Hosts of Dracunculus medinensis, Chad
- Time Lags between Exanthematous Illness Attributed to Zika Virus, Guillain-Barré Syndrome, and Microcephaly, Salvador, Brazil
- Use of Unamplified RNA/cDNA-Hybrid Nanopore Sequencing for Rapid Detection and Characterization of RNA Viruses
- Importation of Hybrid Human-Associated *Trypanosoma cruzi* Strains of Southern South American Origin, Colombia
- · Lyssavirus in Indian Flying Foxes, Sri Lanka
- Survival and Growth of *Orientia tsutsugamushi* in Conventional Hemocultures
- Chagas Disease Screening in Maternal Donors of Publicly Banked Umbilical Cord Blood, United States
- Multilocus Sequence Typing Tool for *Cyclospora* cayetanensis

To revisit the February 2016 issue, go to: https://wwwnc.cdc.gov/eid/articles/issue/22/2/table-of-contents

Improving Quality of Patient Data for Treatment of Multidrug- or Rifampin-Resistant Tuberculosis

Appendix

Data Dictionary for MDR/RR TB IPD

The tables within this section pertain to the data elements optimally preferred for collection during the conduct of observational studies or in routinely collected programmatic data, along with their requested coding to ensure uniformity across studies. Caveats and additional information on specific elements are contained within the main text of the online report.

Facility Information						
Field	Variable	Additional Information	Format	Category Coding	Category Labeling	
COUNTRY	Country	Country of the primary source	Char		_	
TREATING_SITE	Treating Site Name	Name of the primary source	Char			
SITE_ID	Treating Site Identifier	Site ID number	Char			

Patient Identifier and Demographics						
Field	Variable	Additional Information	Format	Category Coding	Category Labeling	
PATIENT_ID	Patient Identifier	Patient ID number in country database	Char			
YEAR	Year	Year of treatment start for this episode	Num ###			
AGE	Age	Age of the patient in years	Num ###			
			Category	F	Female	
SEX	Sex	Patient's biologic sex at birth		M	Male	
				U	Unknown	
WEIGHT	Weight	Patient's weight in kilograms	Num ###			
HEIGHT	Height	Patient's height in centimeters	Num ###			
ВМІ	Body Mass Index	Patient's body mass index in kilograms per meters-squared	Num ###			

Patient Baseline Characteristics						
Field	Variable	Additional Information	Format	Category Coding	Category Labeling	
SMOKINGSTATUS	Smoking Status	The patient's smoking status at start of treatment	Category	Current Ex Never	Current Smoker Ex-Smoker Never Smoker Unknown	
SMOKINGPACKPERDAY	Packs Smoked Per Day	Total number of packs per day smoked at start of treatment (if current smoker)	Num ###	U	Officiown	
SMOKINGTOTALPACKYEAR	Total Pack Years	Total number of pack years smoked (if current- or ex-smoker)	Num ###			
ALCOHOL	Alcohol Use	Does the patient drink (defined as ≥1 drink per week in men or women)	Category	Y N U	Yes No Unknown	
ALCOHOLABUSE	Alcohol Abuse Disorder	If the patient drinks, do they meet the definition of alcohol abuse (≥14 drinks per week in men or ≥7 drinks per week in women)	Category	Y N U	Yes No Unknown	
DM	Diabetes Mellitus	Is the patient diagnosed with diabetes?	Category	Y N U	Yes No Unknown	
INSULINDEPENDENT	Type 1 Diabetes Mellitus	Is the patient insulin dependent (if having diabetes)?	Category	Y N	Yes No Unknown	
HBA1C	Hemoglobin A1c Level	Patients HbA1c measure defined in percent (%)	Num ###		• • • • • • • • • • • • • • • • • • • •	
RENALFAILURE	Presence of Renal Failure	Does the patient have renal failure?	Category	Y N U	Yes No Unknown	
НЕРВ	Hepatitis B	Does the patient have hepatitis B?	Category	Y N U	Yes No Unknown	
HEPC	Hepatitis C	Does the patient have hepatitis C?	Category	Y N U	Yes No Unknown	
OTHERLIVER	Other Liver Condition	Does the patient have liver conditions other than hepatitis B or hepatitis C?	Category	Y N U	Yes No Unknown	
HIV	HIV	What is the patient's HIV status?	Category	Pos Neg	Positive Negative Unknown	
HIV_DIAGNOSISYEAR	Year HIV Diagnosed	If the patient is HIV-positive, the year HIV was diagnosed	Num ###		CHAIGHT	
CD4	CD4 Count	If the patient is HIV-positive, what is their CD4 count at treatment start (cells/µL)?	Num ###			
VIRALLOAD	Viral Load	If the patient is HIV-positive, what is their viral load at treatment start (copies/ml)	Num ###			
ART	Use of Antiretroviral Treatment	If the patient is HIV-positive, are they on antiretroviral treatment?	Category	Y N U	Yes No Unknown	
ART_STARTYEAR	Year Antiretroviral Treatment Started	If the patient is on antiretroviral treatment, what year did they start?	Num ###			
ART_REGIMEN	Antiretroviral Treatment Regimen	What is the antiretroviral treatment regimen? List each drug, separated by a comma, using the provided abbreviations with this dictionary.	Char			

PASTTX Previous Treatment Previous Treatment Previous Treatment with First-Line Drugs Treatment with First-Line Drugs Treatment with Second-Line Drugs Siven for -30 d? Pervious Treatment with Second-Line Previous Treatment		ı	Previous Treatment Information			
Treatment tuberculosis treatment for >30 d? Previous Treatment with First-Line Drugs for >30 d? RECEIVEDSLD RECEIVEDSLD Previous Treatment with First-Line Drugs for >30 d? Previous Treatment with First-Line drugs given for >30 d? Previous Treatment with Second-Line Drugs Year of Most Recent Previous Treatment OutDPASTTX1* Part of Second-Most Recent Previous Treatment Year of Second-Most Recent Previous Treatment The drug-regimen given to the patient at the end of their second-most recent previous tuberculosis treatment to the patient during the second-most recent previous tuberculosis treatment outcome REGIMENPASTTX2* Part of Second-Most Recent Previous Treatment Previous Treatment The drug-regimen given to the patient received previous tuberculosis treatment poisode. The drug-regimen given to the patient during the second-most recent previous tuberculosis treatment poisode. The drug-regimen given to the patient during the second-most recent previous tuberculosis treatment outcome for Second-Most Recent Previous Treatment Outcome for Second-Most Recent Previous Treatment Treatment Previous Treatment Treatment Previous Treatment Treatment Previous Treatment Treatment Treatment Treatment Toutcome for Second-Most Recent Previous Treatment Treatment Treatment Treatment Treatment Toutcome for Second-Most Recent Previous Treatment Treatment Treatment Treatment Treatment Treatment Toutcome for Second-Most Recent Previous Treatment Treatment Treatment Treatment Treatment Toutcome for Second-Most Recent Previous Treatment Treatment Toutcome Toutcome for Second-Most Recent Previous Treatment Treatment Toutcome Toutcome Toutcome Toutcome	Field	Variable	Additional Information	Format		
Previous Treatment with First-Line Drugs RECEIVEDSLD RECEIVE STANDARY RECEIV	PASTTY	Previous		Category		
RECEIVEDFLD Previous Treatment with First-Line Drugs for >30 d? Previous Treatment with Second-Line Drugs Previous Treatment with Second-Line Drugs Year of Most Recent Previous Treatment OUTPASTTX2* Pagimen Used for Second-Most Recent Previous Treatment REGIMENPASTTX2* Pagimen Used for Second-Most Recent Previous Treatment Previous Treatment Previous Treatment End-of-Treatment Uberculosis treatment outcome Previous Treatment Previous Treatment Previous Treatment Previous Treatment End-of-Treatment Uberculosis treatment. Previous Treatment Previous Treatment Previous Treatment during the second-most recent previous tuberculosis treatment. Previous Treatment Previous Treatment Previous Treatment Previous Treatment Dutcome for Second-Most Recent Previous Treatment Failure Previous Treatment Previous Treatment Previous Treatment Previous Treatment Previous Treatment Previous Treatment Previous Treatment Previous Treatment Previous T	FASTIA	Treatment		Category	N	No
First-Line Drugs for 30 d? Previous Treatment with Second-Line Drugs (1) If the patient has received previous tuberculosis treatment, was treatment with second-Line drugs given for >30 d? Year of Most Recent Previous Treatment Regimen Used for Most Recent Previous Treatment OUTPASTTX1* Pare of Second-Most Recent Previous Treatment Year of Second-Most Recent Previous Treatment Previous Treatment Previous Treatment Outcome for Most Recent Previous tuberculosis treatment. Find dug-regimen given to the patient at the end of their most recent previous tuberculosis treatment for their second-most recent treatment episode. The quartened previous tuberculosis treatment for their second-most recent treatment episode. The drug-regimen given to the patient at the end of their second-most recent treatment for their second-most recent treatment episode. The drug-regimen given to the patient at the end of their second-most recent treatment episode. The drug-regimen given to the patient during the second-most recent previous tuberculosis treatment. Char Category N No Category N No No Category Char Category Category N No No Category N No No Category N No Category N No Num ### Char Category Char Ca	RECEIVEDELD		tuberculosis treatment, was	Category	Υ	Yes
Treatment with Second-Line Drugs given for 3-30 d? Year of Most Recent Previous Treatment OUTPASTTX1* Pear of Second-Line Drugs given for 3-30 d? OUTPASTTX2* REGIMENPASTTX2* REGIMENPASTTX3* REGIMENPASTTX3* REGIMENPASTTX3* REGIMENPASTTX3* REGIMENPASTTX4* REGIMENPASTTX4* REGIMENPASTTX4* REGIMENPASTTX4* REGIMENPASTTX4* REGIMENPASTTX4	TREGETYEST ES		for >30 d?	outogory	N	No
YEARPASTTX1* Second-Line prugs given for 30 d? Year of Most Recent Previous Treatment Regimen Used for Most Recent Previous Treatment OUTPASTTX1* Parament The drug-regimen given to the patient at the end of their recorded for the patient received previous tuberculosis treatment The end-of-treatment outcome recorded for the patient at the end of their most recent previous tuberculosis treatment The quartenet outcome recorded for the patient at the end of their most recent previous tuberculosis treatment outcome Year of Second-Most Recent Previous Treatment Year of Second-Most Recent Previous Treatment The year the patient received previous tuberculosis treatment for their second-most recent previous tuberculosis treatment The drug-regimen given to the patient at the end of their most recent previous tuberculosis treatment for their second-most recent treatment episode. The year the patient received previous tuberculosis treatment for their second-most recent previous tuberculosis treatment. List each drug, separated by a comma, using the provided abbreviations with this dictionary. End-of-Treatment OutpastTX2* DUTPASTTX2* End-of-Treatment OutpastTX2* End-of-Treatment OutpastTX2* The drug-regimen given to the patient at the end of their second-most recent previous tuberculosis treatment. List each drug, separated by a comma, using the provided abbreviations with this dictionary. The drug-regimen given to the patient at the end of their second-most recent previous tuberculosis treatment. List each drug, separated by a comma, using the provided abbreviations with this dictionary. The drug-regimen given to the patient at the end of their second-most recent previous tuberculosis treatment. List each drug, separated by a comma, using the provided abbreviations with this dictionary. End-of-Treatment OutpastTX2* The drug-regimen given to the patient at the end of their second-most recent previous tuberculosis treatment to the patient at the end of their second-most recent previous tuberculosis trea	DECEIVEDSI D	Treatment with tuberculosis treatment, was	Cotogony	Υ	Yes	
REGIMENPASTTX1* Recent Previous Treatment REGIMENPASTTX1* Regimen Used for Most Recent Previous tuberculosis treatment. List each drug, separated by a comma, using the provided abbreviations with this dictionary. Part of Second-Most Recent Previous Treatment Previous Treatment Year of Second-Most Recent Previous Treatment REGIMENPASTTX2* REGIMENPASTTX4* REGIMENPASTTX4*	RECEIVEDSLD		given for >30 d?	Category	N	No
REGIMENPASTTX1* Regimen Used for Most Recent Previous Treatment Previous Treatment Regimen Used for Most Recent Previous tuberculosis treatment. List each drug, separated by a comma, using the provided abbreviations with this dictionary. End-of-Treatment Outcome for Most Recent Previous Treatment Year of Second-Most Recent Previous Treatment Previous Treatment Year of Second-Most Recent Previous Treatment The year the patient received previous tuberculosis treatment of their second-most recent treatment episode. Regimen Used for Second-Most Recent Previous Treatment The drug-regimen given to the patient during the second-most recent previous tuberculosis treatment. List each drug, separated by a comma, using the provided abbreviations with this dictionary. End-of-Treatment Outcome for Second-Most Recent Previous Treatment The end-of-treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. The end-of-treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. End-of-Treatment The end-of-treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. End-of-Treatment The end-of-treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. End-of-Treatment The end-of-treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. End-of-Treatment The end-of-treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. End-of-Treatment The end-of-treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. E	YEARPASTTX1*	Recent Previous	received previous tuberculosis	Num ###		
OUTPASTTX1* End-of-Treatment Outcome for Most Recent Previous Treatment Outcome for Most Recent Previous Treatment Year of Second-Most Regimen Used for Second-Most Recent Previous Treatment REGIMENPASTTX2* OUTPASTTX2* End-of-Treatment OUTPASTTX2* End-of-Treatment OUTPASTTX2* End-of-Treatment OUTPASTTX2* End-of-Treatment OUTPASTTX2* Duriphical Completed Treatment Failure Lost Lost to Follow-up U Unknown Num ### Regimen Used for Second-Most Recent Previous Treatment The drug-regimen given to the patient during the second-most recent previous tuberculosis treatment. List each drug, separated by a comma, using the provided abbreviations with this dictionary. End-of-Treatment Outcome for Second-Most Recent Previous Treatment The end-of-treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. End-of-Treatment Outcome for Second-Most Recent Previous Treatment The end-of-treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. Category Complete Completed Treatment Characteristics The end-of-treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. Category Lost Completed Treatment Characteristics The end-of-treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. Category Lost Completed Treatment Completed Treatment The end-of-treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. Category	REGIMENPASTTX1*	for Most Recent Previous	patient during the most recent previous tuberculosis treatment. List each drug, separated by a comma, using the provided	Char		
Outcome for Most Recent Previous Treatment Year of Second-Most Recent Previous Treatment Regimen Used for Second-Most Recent Previous Treatment Outpastive Regimen Used for Second-Most Recent Previous Treatment Num ### Category The year the patient received previous tuberculosis treatment for their second-most recent treatment episode. The drug-regimen given to the patient during the second-most recent previous tuberculosis treatment. List each drug, separated by a comma, using the provided abbreviations with this dictionary. End-of-Treatment Outcome for Second-Most Recent Previous Treatment Previous Treatment Their second-most recent previous tuberculosis treatment at the end of their second-most recent previous tuberculosis treatment. Char Cure Completed Treatment Completed Treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. Category Unknown				Category	Cure	
Treatment tuberculosis treatment. Year of Second- Most Recent Previous Treatment tuberculosis treatment tuberculosis treatment tuberculosis treatment for their second-most recent treatment episode. Regimen Used for Second-Most Recent Previous Treatment tuberculosis treatment tuberculosis treatment tuberculosis treatment. End-of-Treatment Outcome for Second-Most Recent Previous Treatment Tuberculosis treatment outcome for Second-Most Recent Previous Treatment Tuberculosis treatment. End-of-Treatment Outcome for Second-most recent previous tuberculosis treatment outcome for their second-most recent previous tuberculosis treatment outcome for their second-most recent previous tuberculosis treatment. End-of-Treatment Outcome for Second-Most Recent Previous Treatment Tuberculosis treatment outcome for their second-most recent previous tuberculosis treatment tuberculosis treatment. End-of-Treatment Outcome for Second-Most Recent Previous Treatment Tuberculosis treatment tuberculosis treatment. End-of-Treatment Outcome for Second-Most Recent Previous Tuberculosis treatment tuberculosis treatment. End-of-Treatment Outcome for Second-Most Recent Previous Tuberculosis treatment tuberculosis treatment. End-of-Treatment Outcome for Second-Most Recent Previous Tuberculosis treatment tuberculosis treatment. End-of-Treatment Outcome for Second-Most Recent Previous Tuberculosis treatment tuberculosis treatment. End-of-Treatment Outcome for Second-Most Recent Previous Tuberculosis treatment Tuberculosis treat	OLITO A OTT VA *	Outcome for Most	recorded for the patient at the end of their most recent previous		Complete	
YEARPASTTX2* Year of Second-Most Recent Previous Treatment OUTPASTTX2* Pear of Second-Most Recent Previous Treatment Outpast Tx2* Pear of Second-Most Recent Previous Treatment Previous Treatment Regimen Used for Second-Most Recent Previous Treatment Outpast Tx2* End-of-Treatment Outcome for Second-Most Recent Previous Treatment Outpast Tx2* OUTPASTTX2* The year the patient received previous tuberculosis treatment on the patient during the second-most recent previous tuberculosis treatment. Char Char Cure Complete Complete Complete Treatment The end-of-treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. Category U Unknown	OUTPASTIXI				Fail	
YEARPASTTX2* Year of Second-Most Recent Previous tuberculosis treatment of their second-most recent treatment episode. The drug-regimen given to the patient during the second-most recent previous tuberculosis treatment. End-of-Treatment Outcome for Second-Most Recent Previous Treatment OUTPASTTX2* Pyear of Second-Most Recent Previous tuberculosis treatment outloome recent previous with this dictionary. End-of-Treatment Outcome for Second-Most Recent Previous Treatment Outcome for Second-most recent previous tuberculosis treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. Char Char Cure Cure Complete Treatment Category Fail Treatment Failure Lost Lost to Follow-up U Unknown		Treatment			Lost	Lost to Follow-up
Most Recent Previous Treatment Regimen Used for Second-Most Recent Previous Treatment OUTPASTTX2* Most Recent Previous Treatment Previous Treatment Regimen Used for Second-Most Recent Previous Treatment Outcome for Second-most recent previous tuberculosis treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. Category Cure Complete Treatment Category Fail Treatment Failure Lost Lost to Follow-up U Unknown					U	Unknown
REGIMENPASTTX2* Regimen Used for Second-Most Recent Previous Treatment Patient during the second-most recent previous tuberculosis treatment. List each drug, separated by a comma, using the provided abbreviations with this dictionary. End-of-Treatment Outcome for Second-Most Recent Previous Treatment Outrain The end-of-treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. Char Char Cure Complete Complete Treatment Treatment Fail Treatment Failure Lost Lost to Follow-up U Unknown	YEARPASTTX2*	Most Recent Previous	previous tuberculosis treatment for their second-most recent treatment episode.	Num ###		
OUTPASTTX2* End-of-Treatment Outcome for Second-Most Recent Previous Treatment The end-of-treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. The end-of-treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. The end-of-treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. Category Fail Treatment Failure Lost Lost to Follow-up U Unknown	REGIMENPASTTX2*	for Second-Most Recent Previous	patient during the second-most recent previous tuberculosis treatment. List each drug, separated by a comma, using the provided	Char		
Outcome for Second-Most Recent Previous Treatment Outcome for Second-Most Recent Previous Treatment Outcome for Second-Most Recent Previous Treatment The end-of-treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. The end-of-treatment outcome recorded for the patient at the end of their second-most recent previous tuberculosis treatment. Category Fail Treatment Failure Lost Lost to Follow-up U Unknown	OLITDA STTV2*	End-of-Treatment			Cure	
Recent Previous Treatment their second-most recent previous tuberculosis treatment. Their second-most recent previous tuberculosis treatment. Fail Treatment Failure Lost Lost to Follow-up U Unknown		Outcome for	recorded for the patient at the end of their second-most recent previous	Category		Treatment
Treatment tuberculosis treatment. Lost Lost to Follow-up U Unknown	00117011772					
U Unknown			tuberculosis treatment.			
					U	Unknown

		Disease Characteristics			
Field	Variable	Additional Information	Format	Category Coding	Category Labeling
	Site of			PTB	Pulmonary TB
DISEASE_SITE	Tuberculosis	The site of tuberculosis disease	Category	EPTB	Extrapulmonary TB
	Disease	diagnosed in the patient		Both	Both
				Miliary	Miliary TB
				Genital	Genitourinary TB
				CNS	Central Nervous System TB
	Delas and Oite of			Periton	TB Peritonitis
EVEDADUIM SITE	Primary Site of	If extrapulmonary tuberculosis is	Cotogoni	Pericar	TB Pericarditis
EXTRAPULM_SITE	Extrapulmonary Tuberculosis	diagnosed, the primary site affected	Category	Lymph	TB Lymphadenitis
	Tuberculosis			Pleural	Pleural TB
				GI	Gastrointestinal TB
				Bone	Bone TB
				Joint	Joint TB
				Other	Other
CAVITATION_BASE*	Lung Cavitation	Was there presence of lung		Υ	Yes
		cavitation on chest x-ray at treatment start?	Category	N	No
				U	Unknown
		Was there presence of bilateral disease on chest X-ray at treatment start?		Υ	Yes
BILATERAL_BASE*	Bilateral Disease		Category	N	No
				U	Unknown
		What was the patient's acid-fast		Pos	Positive
		bacilli smear result (taken ≤1 mo		Neg	Negative
	Acid-Fast Bacilli	after treatment start)?		Contam	Contaminated
AFB_BASE	Smear Result	Consider all samples taken over this time frame and consider positive if any were positive (i.e., scanty or greater).	Category	ND	Not Done
		What was the patient's sputum		Pos	Positive
		culture result (taken <1 mo after		Neg	Negative
CULTURE BASE	Sputum Culture	treatment start)?	Category	Contam	Contaminated
OSET SINE_BAGE	Result	Consider all samples taken over this frame and consider positive if any were positive.	Calogory	ND	Not Done
CHITUDEMEDIA	Culture Media	If culture was done, what media was	0-4	Solid	Solid Media
CULTUREMEDIA	Used	used for the result reported?	Category	Liquid	Liquid Media
*Baseline refers to any evidence	of cavitation or bilatera	al disease within 30 d of treatment start.		-	

Genotypic DST						
Variable	Additional Information	Format	Category Coding	Category Labeling		
Genotypic DST	Were genotypic DST	Category	Y	Yes No		
USEU						
Gene Xpert Used		Category		Yes No		
Date of Gene Xpert	Date of Gene Xpert used for diagnosis <mm dd="" yy=""></mm>	Date		110		
Gene Xpert MTB	What was the result for MTB on Gene	Category	Pos Neg	Positive Negative		
	Xpert?			Contaminated		
Rifampin	What was the result	Catagory	R S	Resistant Susceptible		
Resistance Result	on Gene Xpert?	Category	Contam	Contaminated		
First Line LDA	Was first-line LPA		Υ	Yes		
Used	used after TB diagnosis?	Category	N	No		
Date of First-Line LPA	Date of first-line LPA used after TB diagnosis <mm dd="" yy=""></mm>	Date				
First-Line LPA MTB Result	What was the result for MTB on first-line LPA?	Category	Pos	Positive		
			Neg	Negative		
				Contaminated		
Isoniazid Resistance	for isoniazid resistance on first-line			Resistant		
		Category	Contam	Susceptible Contaminated		
			R	Resistant		
		Category		Susceptible		
Resistance	on first-line LPA?		Contam	Contaminated		
	Was second-line LPA		Υ	Yes		
Second-Line LPA Used	performed after TB diagnosis?	Category	N	No		
Date of Second- Line LPA	Date of second-line LPA used after TB diagnosis <mm dd="" yy=""></mm>	Date				
Second-Line I DA	What was the result		Pos	Positive		
		Category	Neg	Negative		
	line LPA?		Contam	Contaminated		
	What was the result			Resistant		
	for second-line	0-4-	S	Susceptible		
Resistance Result	injectable resistance on second-line LPA?	Category	Contam	Contaminated		
Second-Line LPA	What was the result		R	Resistant		
Fluoroquinolone	for fluoroquinolone	Category	S	Susceptible		
Resistance Result	resistance on second- line LPA?		Contam	Contaminated		
	Genotypic DST Used Gene Xpert Used Date of Gene Xpert Gene Xpert MTB Result Gene Xpert Rifampin Resistance Result First-Line LPA Used Date of First-Line LPA First-Line LPA Isoniazid Resistance Result First-Line LPA Isoniazid Resistance Result First-Line LPA Isoniazid Resistance Result First-Line LPA Second-Line LPA Used Date of Second-Line LPA Second-Line LPA MTB Result Second-Line LPA MTB Result Second-Line LPA MTB Result Second-Line LPA Second-Line LPA Second-Line LPA Second-Line Ipectable Resistance Result Second-Line LPA Resistance Result	Genotypic DST Used Were genotypic DST techniques used? Gene Xpert Used Date of Gene Xpert used for diagnosis? Gene Xpert MTB Result What was the result for rifampin resistance on Gene Xpert? First-Line LPA Used Seristance Pasult Seristance Result PA? First-Line LPA Used Second-Line LPA Used Second-Line LPA Used Second-Line LPA Second-Line LPA Second-Line LPA Second-Line LPA Sesistance Result Second-Line LPA Second-Line LPA? Second-Line LPA? Second-Line LPA Second-Line LPA Second-Line LPA? Second-Line LPA Second-Line LPA Second-Line LPA? Second-Line LPA Second-Line LPA?	Variable Additional Information Format Genotypic DST Used Were genotypic DST techniques used? Category Gene Xpert Used Was Gene Xpert used for diagnosis? Category Date of Gene Xpert used for diagnosis <mm dd="" yy=""> Date of Gene Xpert used for diagnosis <mm dd="" yy=""> Date of Gene Xpert used for diagnosis <mm dd="" yy=""> Category Gene Xpert MTB Result What was the result for iffampin resistance on Gene Xpert? Category Gene Xpert Rifampin Resistance Result Was first-line LPA used after TB diagnosis? Category Date of First-Line LPA Used Was first-line LPA used after TB diagnosis <mm d="" yy=""> Category First-Line LPA MTB Result What was the result for MTB on first-line LPA? Category First-Line LPA Rifampin Resistance Result What was the result for isoniazid resistance on first-line LPA? Category First-Line LPA Used What was the result for iffampin resistance on first-line LPA? Category Second-Line LPA Used What was the result for iffampin resistance on first-line LPA? Category Second-Line LPA Used What was the result for iffampin resistance on first-line LPA? Category Second-Line LPA MTB Result What was the result for MTB on second</mm></mm></mm></mm>	Variable		

*Baseline DST refers to any sample taken within 90 d of treatment start, up to 30 d after treatment start. Every effort should be made to have reliable DST results; if genotypic tests are not used, phenotypic tests should be performed. If genotypic techniques for detection other than those listed in this table are in use (e.g., pncA for pyrazinamide), they may be appended to this section in a similar format (e.g., Test Done, Date of Test, Results of Test).

Phenotypic DST						
Field*	Variable	Additional Information	Format	Category Coding	Category Labeling	
PHENODST	Phenotypic DST Done	Was phenotypic DST performed?	Category	Y N	Yes No	
DATE_PHENODST	Date of Phenotypic DST	Date of phenotypic DST done after TB diagnosis <mm dd="" yy=""></mm>	Date	IN	NO	
	,	What was the result for isoniazid		R	Resistant	
DST_H_BASE	Isoniazid Resistance	resistance (MIC >0.1–0.2 µg/ml on	Category	S	Susceptible	
DOT_IT_BASE	Result	MGIT) on phenotypic DST?	Category	Contam	Contaminated	
		merry on phonotypic 201.		ND	Not Done	
	High-Level	What was the result for high-level		R	Resistant	
DST HIGHH BASE	Isoniazid	isoniazid resistance (MIC >1-2 µg/ml	Category	S	Susceptible	
	Resistance	on MGIT) on phenotypic DST?	3. 7	Contam	Contaminated	
	Result	, , ,		ND	Not Done	
	Rifampin	NAME of the second formal formal and		R	Resistant	
DST_R_BASE	Resistance	What was the result for rifampin	Category	S Contam	Susceptible	
	Result	resistance on phenotypic DST?		ND	Contaminated Not Done	
				R	Resistant	
	Ethambutol	What was the result for ethambutol		S	Susceptible	
DST_E_BASE	Resistance		Category	Contam	Contaminated	
	Result resistance on phenotypic DST?		ND	Not Done		
				R	Resistant	
	Pyrazinamide	What was the result for pyrazinamide		S	Susceptible	
DST_Z_BASE	Resistance	resistance on phenotypic DST?	Category	Contam	Contaminated	
	Result	resistance on pnenotypic DST?	= *	ND	Not Done	
				R	Resistant	
	Amikacin	What was the result for amikacin resistance on phenotypic DST?		S	Susceptible	
DST_AM_BASE	Resistance		Category	Contam	Contaminated	
	Result			ND	Not Done	
		What was the result for kanamycin resistance on phenotypic DST?		R	Resistant	
	Kanamycin			S	Susceptible	
DST_KM_BASE	Resistance		Category	Contam	Contaminated	
	Result			ND	Not Done	
	Ci-	What was the result for capreomycin	Category	R	Resistant	
DOT OM BASE	Capreomycin Resistance			S	Susceptible	
DST_CM_BASE	Resistance	resistance on phenotypic DST?		Contam	Contaminated	
	Nesuit	. ,,		ND	Not Done	
	Ofloyooin	What was the result for ofloxacin resistance on phenotypic DST?	Category	R	Resistant	
DST_OFX_BASE	Ofloxacin Resistance			S	Susceptible	
DOT_OFX_BASE	Result			Contam	Contaminated	
	rtoodit			ND	Not Done	
	Ciprofloxacin			R	Resistant	
DST_CFX_BASE	Resistance	What was the result for ciprofloxacin	Category	S	Susceptible	
200.7_27.02	Result	resistance on phenotypic DST?	outogo.)	Contam	Contaminated	
				ND	Not Done	
	Moxifloxacin	100		R	Resistant	
DST_MFX_BASE	Resistance	What was the result for moxifloxacin	Category	S	Susceptible	
	Result	resistance on phenotypic DST?		Contam ND	Contaminated Not Done	
	Levofloxacin	What was the result for levofloxacin		R S	Resistant Susceptible	
DST_LFX_BASE	Resistance	resistance on phenotypic DST?	Category	Contam	Contaminated	
	Result	resistance on phenotypic bor :		ND	Not Done	
				R	Resistant	
	Streptomycin	What was the result for streptomycin		S	Susceptible	
DST_S_BASE	Resistance	resistance on phenotypic DST?	Category	Contam	Contaminated	
	Result	process, pro Bot .		ND	Not Done	
				R	Resistant	
	Ethionamide	What was the result for ethionamide	Category	S	Susceptible	
DST_ETO_BASE	Resistance	resistance on phenotypic DST?		Contam	Contaminated	
	Result	, , , , , ,		ND	Not Done	
	Prothionamide	What was the result for		R	Resistant	
DST_PTO_BASE	Resistance	prothionamide resistance on	Category	S	Susceptible	
	Result	phenotypic DST?	,	Contam	Contaminated	
	Nesull	prienotypic DOT:				

		Phenotypic DST			
Field*	Variable	Additional Information	Format	Category Coding	Category Labeling
				ND	Not Done
	Ouglas a min a			R	Resistant
DST CS BASE	Cycloserine Resistance	What was the result for cycloserine	Cotogomi	S	Susceptible
DS1_CS_BASE	Result	resistance on phenotypic DST?	Category	Contam	Contaminated
	Result			ND	Not Done
	T = 2'=1'd = 2 =			R	Resistant
DOT TOD DACE	Terizidone	What was the result for terizidone	0-4	S	Susceptible
DST_TRD_BASE	Resistance Result	resistance on phenotypic DST?	Category	Contam	Contaminated
	Result			ND	Not Done
DST_PAS_BASE	Para-Amino-			R	Resistant
	Salicylic Acid		Category	S	Susceptible
	Resistance			Contam	Contaminated
	Result	phenotypic DST?		ND	Not Done
	1. 1. 1. 1	What was the result for linezolid resistance on phenotypic DST?		R	Resistant
DOT 17D DAGE	Linezolid		Category	S	Susceptible
DST_LZD_BASE	Resistance Result			Contam	Contaminated
	Result			ND	Not Done
	01 () ;		0 1	R	Resistant
DOT OFT DAGE	Clofazimine	What was the result for clofazimine		S	Susceptible
DST_CFZ_BASE	Resistance	resistance on phenotypic DST?	Category	Contam	Contaminated
	Result			ND	Not Done
	D 1 11			R	Resistant
DOT DDO DAOE	Bedaquiline	What was the result for bedaquiline	0-1	S	Susceptible
DST_BDQ_BASE	Resistance Result	resistance on phenotypic DST?	Category	Contam	Contaminated
	Result			ND	Not Done
	Dalamanid			R	Resistant
DOT DIM DACE	Delamanid	What was the result for delamanid	Category	S	Susceptible
DST_DLM_BASE	Resistance	resistance on phenotypic DST?		Contam	Contaminated
	Result			ND	Not Done

*Baseline DST refers to any sample taken within 90 d of treatment start, up to 30 d after treatment start. Additional drugs for which phenotypic DST is available can be reported (e.g., Pretomanid). Within all shared data, the method and critical concentration used for each drug must be recorded.

Follow-Up DST and Acquired Drug Resistance					
Field	Variable	Additional Information	Format	Category Coding	Category Labeling
FOLLOWUP_DST	Follow-up DST Performed	Was there follow-up DST performed?	Category	Y N	Yes No
FOLLOWUPDST1_DATE*	Date of First Follow-up DST	Date of first follow-up DST <mm dd="" yy=""></mm>	Date		
FOLLOWUPDST_RES1	Resistant Isolates on First Follow-up DST	List newly discovered resistances not found on baseline DST, due to missingness or baseline susceptibility. If none discovered, list "no change in DST." List each drug, separated by a comma, using the provided abbreviations with this dictionary.	Char		
FOLLOWUPDST_SUS1	Susceptible Isolates on First Follow-up DST	List newly discovered susceptible drugs not found on baseline DST, due to missingness or baseline resistance. If none discovered, list "no change in DST." List each drug, separated by a comma, using the provided abbreviations with this dictionary.	Char		
ACQUIRED_RESISTANCE†	Acquired Drug Resistance	List the drugs that the strain was shown to acquire resistance to during any follow-up DST (defined as previously identified susceptibility and subsequent resistance on follow-up DST). List each drug, separated by a comma, using the provided abbreviations with this dictionary.	Char		

^{*}Additional follow-up DST results can be entered following a similar format.

†Acquired resistance can be reported in a separate row but is not necessary as it can be calculated by the data analyst with the above collected data.

Regimen Information*								
Field	Variable	Additional Information	Format	Category Coding	Category Labeling			
STARTINGREGIMENTYPE	Regimen Type at Start of	List the starting regimen type: short (intended duration ≤12 mo) or long	Category	Short	Short Regimen			
	Treatment	(intended duration ≥18 mo) Date of second-line drug initiation in		Long	Long Regimen			
TXSTART_DATE	Treatment Start Date	this treatment episode ">mm/dd/yy>	his treatment episode Date Cmm/dd/yy>					
INITIAL_REGIMEN	Starting Treatment Regimen	List the drugs the patient is on at the start of treatment. List each drug, separated by a comma, using the provided abbreviations with this dictionary.	Char					
H_START	Isoniazid Start Date	Date standard-dose isoniazid was introduced into the patient's regimen. <mm dd="" yy=""></mm>	Date					
H_STOP	Isoniazid End Date	Date standard-dose isoniazid was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date					
HIGHH_START	High-Dose Isoniazid Start Date	Date high-dose isoniazid was introduced into the patient's regimen. <mm dd="" yy=""></mm>	Date					
HIGHH_STOP	High-Dose Isoniazid End Date	Date high-dose isoniazid was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date					
E_START	Ethambutol Start Date	Date ethambutol was introduced into the patient's regimen. <pre><mm dd="" yy=""></mm></pre>	Date					
E_STOP	Ethambutol End Date	Date ethambutol was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date					
Z_START	Pyrazinamide Start Date	Date pyrazinamide was introduced into the patient's regimen. <mm dd="" yy=""></mm>	Date					
Z_STOP	Pyrazinamide End Date	Date pyrazinamide was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date					
S_START	Streptomycin Start Date	Date streptomycin isoniazid was introduced into the patient's regimen. <mm dd="" yy=""></mm>	Date					
S_STOP	Streptomycin End Date	Date streptomycin was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date					
RFB_START	Rifabutin Start Date	Date rifabutin was introduced into the patient's regimen. <mm dd="" yy=""></mm>	Date					
RFB_STOP	Rifabutin End Date	Date rifabutin was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date					
AM_START	Amikacin Start Date	Date amikacin was introduced into the patient's regimen. <mm dd="" yy=""></mm>	Date					
AM_STOP	Amikacin End Date	Date amikacin was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date					
KM_START	Kanamycin Start Date	Date kanamycin was introduced into the patient's regimen. <mm dd="" yy=""></mm>	Date					
KM_STOP	Kanamycin End Date	Date kanamycin was permanently removed from the patient's regimen	Date					

Regimen Information*							
Field	Variable	Additional Information	Format	Category Coding	Category Labeling		
		<mm dd="" yy=""></mm>					
CM_START	Capreomycin Start Date	Date capreomycin was introduced into the patient's regimen. <mm dd="" yy=""></mm>	Date				
CM_STOP	Capreomycin End Date	Date capreomycin was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date				
OFX_START	Ofloxacin Start Date	Date ofloxacin was introduced into the patient's regimen. <pre><mm dd="" yy=""></mm></pre>	Date				
OFX_STOP	Ofloxacin End Date	Date ofloxacin was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date				
CFX_START	Ciprofloxacin Start Date	Date ciprofloxacin was introduced into the patient's regimen. <pre><mm dd="" yy=""></mm></pre>	Date				
CFX_STOP	Ciprofloxacin End Date	Date ciprofloxacin was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date				
MFX_START	Moxifloxacin Start Date	Date moxifloxacin was introduced into the patient's regimen. <mm dd="" yy=""></mm>	Date				
MFX_STOP	Moxifloxacin End Date	Date moxifloxacin was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date				
LFX_START	Levofloxacin Start Date	Date levofloxacin was introduced into the patient's regimen. <pre><mm dd="" yy=""></mm></pre>	Date				
LFX_STOP	Levofloxacin End Date	Date levofloxacin was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date				
GFX_START	Gatifloxacin Start Date	Date gatifloxacin was introduced into the patient's regimen. <mm dd="" yy=""></mm>	Date				
GFX_STOP	Gatifloxacin End Date	Date gatifloxacin was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date				
SFX_START	Sparfloxacin Start Date	Date sparfloxacin was introduced into the patient's regimen. <mm dd="" yy=""></mm>	Date				
SFX_STOP	Sparfloxacin End Date	Date sparfloxacin was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date				
ETO_START	Ethionamide Start Date	Date ethionamide was introduced into the patient's regimen. <pre><mm dd="" yy=""></mm></pre>	Date				
ETO_STOP	Ethionamide End Date	Date ethionamide was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date				
PTO_START	Prothionamide Start Date	Date prothionamide was introduced into the patient's regimen. <pre><mm dd="" yy=""></mm></pre>	Date				
PTO_STOP	Prothionamide End Date	Date prothionamide was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date				
CS_START	Cycloserine Start Date	Date cycloserine was introduced into the patient's regimen. <pre><mm dd="" yy=""></mm></pre>	Date				
CS_STOP	Cycloserine End Date	Date cycloserine was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date				
TRD_START	Terizidone Start Date	Date terizidone was introduced into the patient's regimen. <pre><mm dd="" yy=""></mm></pre>	Date				

Regimen Information*							
Field	Variable	Additional Information	Format	Category Coding	Category Labeling		
TRD_STOP	Terizidone End Date	Date terizidone was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date				
PAS_START	Para- Aminosalicylic Acid Start Date	Date para-aminosalicylic acid was introduced into the patient's regimen. <mm dd="" yy=""></mm>	Date				
PAS_STOP	Para- Aminosalicylic Acid End Date	Date para-aminosalicylic acid was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date				
LZD_START	Linezolid Start Date	Date linezolid was introduced into the patient's regimen. <mm dd="" yy=""></mm>	Date				
LZD_STOP	Linezolid End Date	Date linezolid was permanently removed from the patient's regimen rmm/dd/yy>	Date				
CFZ_START	Clofazimine Start Date	Date clofazimine was introduced into the patient's regimen. <pre><mm dd="" yy=""></mm></pre>	Date				
CFZ_STOP	Clofazimine End Date	Date clofazimine was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date				
AMXCLV_START	Amoxicillin and Clavulanic Acid Start Date	Date amoxicillin and clavulanic acid was introduced into the patient's regimen. <mm dd="" yy=""></mm>	Date				
AMXCLV_STOP	Amoxicillin and Clavulanic Acid End Date	Date amoxicillin and clavulanic acid was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date				
IPM_START	Imipenem- Cilastatin Start Date	Date imipenem-cilastatin was introduced into the patient's regimen. <mm dd="" yy=""></mm>	Date				
IPM_STOP	Imipenem- Cilastatin End Date	Date imipenem-cilastatin was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date				
MPM_START	Meropenem Start Date	Date meropenem was introduced into the patient's regimen. ">kmm/dd/yy>	Date				
MPM_STOP	Meropenem End Date	Date meropenem was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date				
BDQ_START	Bedaquiline Start Date	Date bedaquiline was introduced into the patient's regimen. <pre><mm dd="" yy=""></mm></pre>	Date				
BDQ_STOP	Bedaquiline End Date	Date bedaquiline was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date				
DLM_START	Delamanid Start Date	Date delamanid was introduced into the patient's regimen. <mm dd="" yy=""></mm>	Date				
DLM_STOP	Delamanid End Date	Date delamanid was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date				
PA_START	Pretomanid Start Date	Date pretomanid was introduced into the patient's regimen. <pre><mm dd="" yy=""></mm></pre>	Date				
PA_STOP	Pretomanid End Date	Date pretomanid was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date				
PCZ_START	Perchlozone Start Date	Date perchlozone was introduced into the patient's regimen. <pre><mm dd="" yy=""></mm></pre>	Date				

Regimen Information*								
Field	Variable	Additional Information	Format	Category Coding	Category Labeling			
PCZ_STOP	Perchlozone End Date	Date perchlozone was permanently removed from the patient's regimen <mm dd="" yy=""></mm>	Date					
TXEND_DATE	Treatment End Date	Date treatment ended in this treatment episode <mm dd="" yy=""></mm>	Date					
	Intended	If the nations started on a short		Υ	Yes			
DURATION_CHANGE	Duration of Regimen Changed	If the patient started on a short regimen, did they switch to a long regimen?	Category	N	No			
CHANGE_DATE	Date of Regimen Duration Change	The date the patient changed from a short regimen to a long regimen <mm dd="" yy=""></mm>	Date					
		What was the reason the regimen		DST	Drug Resistance			
	Reason the	duration changed? This may		NoResp	Non-Response			
CHANGE REASON	Regimen	include: in response to drug	Category	AE	Tolerability			
OTANGE_REAGON	Duration	susceptibility testing, treatment non-	Catogory	Avail	Drug Availability			
	Changed	response, drug availability, drug tolerability, or other. in blank. Stop dates must refer to the date		Other	Other			

^{*}For drugs not used in the regimen, their coding can remain blank. Stop dates must refer to the date that the drug was permanently withdrawn from the regimen. New rows can be added to accommodate drugs not contained in this table.

	Treatment Information								
Field	Variable	Additional Information	Format	Category Coding	Category Labeling				
TXDUR_DAYS	Treatment Duration	Total number of days of treatment, from first to last dose taken	Num ###						
DOT	Directly Observed	Was directly observed therapy used?	Category	Υ	Yes				
201	Therapy	was allestly observed therapy used:	Outogory	N	No				
		State the type of directly observed		Comm	Community				
	Type of Directly	therapy used. Virtual includes methods such as video, mobile text, or medication monitoring, among others.	Category	Hosp	Hospital				
DOT_TYPE	Therapy			Pharm	Pharmacy				
				Virtual	Virtual				
DOT_FREQUENCY	Frequency of DOT Visits	How many days per week is DOT provided to the patient (range 0–7 d)	Num ###						
		What form of patient support was		Employ	Employment				
		provided to patients? This may		Nutri	Nutritional				
SUPPORT	Patient Support	include support from employers (job	Cotogoni	Finance	Financial				
SUFFURI	Provided	security), nutritional support, financial	Category	Other	Other				
		support, or others. If more than one		Multi	Multiple				
		form, please select multiple.		None	None				

	Surgery and Hospitalization Information								
Field	Variable	Additional Information Format		Category Coding	Category Labeling				
	Lung Resection	Did the patient have lung resection		Υ	Yes				
SURGERY	Surgery	surgery related to MDR/RR-TB?	Category	N	No				
	Surgery	Surgery related to MIDIOTOTOTO:		U	Unknown				
				Lobe	Lobectomy				
	Type of Lung	What was the type of lung reception		Pneu	Pneumonectomy				
SURGTYPE	Resection Surgery	What was the type of lung resection surgery?		Wedge	Wedge Resection				
				Other	Other				
				U	Unknown				
SURG_DATE	Date of Surgery	What was the date of surgery?	Date						
		Was the patient hospitalized at any		Υ	Yes				
HOSP	Hospitalization	point during treatment?	Category	N	No				
		point during treatment:		U	Unknown				
HOSPEPISODES	Number of Hospitalization Episodes	What is the total number of hospitalization episodes during treatment?	Num ###						
HOSPDUR_DAYS	Total Hospitalization Duration	What is the total duration of hospitalization during treatment?	Num ###						

		Adverse Event Information		Cotome	Cotogoni
Field*	Variable	Additional Information	Format	Category Coding	Category Labeling
A.F.4	First Adverse	Did the patient experience a serious		SAE	Serious Adverse Event
AE1	Event	adverse event and/or permanently stop the drug	Category	Perm Both	Permanent Stop Both
ΛΓ1 DΛΤΓ	Date of First	What was the date of the permanent	Doto	DOIII	Botti
AE1_DATE	Adverse event	discontinuation of the drug(s)?	discontinuation of the drug(s)?		
AE1_DRUG	Drug Responsible for First Adverse Event	List each drug, separated by a comma, using the provided abbreviations with this dictionary.	Char		
AE1_GRADE	Grade of First Adverse Event	What was the grade of the first adverse event?	Num ###		
AE1_SYSTEMORGAN	System / Organ Class Affected by First Adverse Event	Which system / organ classes were affected by the first adverse event? List each system / organ class, separated by a comma, using the list provided with this dictionary.	Char		
	Outsome of First			Recov	Recovered
AE1_OUTCOME	Outcome of First Adverse Event	What was the outcome of the first adverse event?	Category	NoRecov Died	Not Recovered Died
	Adverse Everit	auverse event:		U	Unknown
	Cooped Adverse	Did the patient experience a serious		SAE	Serious Adverse Event
AE2	Second Adverse Event	adverse event and/or permanently	Category	Perm	Permanent Stop
		stop the drug		Both	Both
AE2_DATE	Date of Second Adverse event	What was the date of the permanent discontinuation of the drug(s)?	Date		
AE2_DRUG	Drug Responsible for Second Adverse Event	List each drug, separated by a comma, using the provided abbreviations with this dictionary.	Char		
AE2_GRADE	Grade of Second Adverse Event	What was the grade of the second adverse event?	Num ###		
AE2_SYSTEMORGAN	System / Organ Class Affected by Second Adverse Event	Which system / organ classes were affected by the second adverse event? List each system / organ class, separated by a comma, using the list provided with this dictionary.	Char		
	Outcome of			Recov	Recovered
AE2_OUTCOME	Second Adverse	What was the outcome of the second adverse event?	Category	NoRecov Died	Not Recovered Died
	Event	adverse event?		U	Unknown
	Third Adverse	Did the patient experience a serious		SAF	Serious Adverse Event
AE3	Event	adverse event and/or permanently	Category	Perm	Permanent Stop
		stop the drug		Both	Both
AE3_DATE	Date of Third Adverse event	What was the date of the permanent discontinuation of the drug(s)?	Date		
AE3_DRUG	Drug Responsible for Third Adverse Event	List each drug, separated by a comma, using the provided abbreviations with this dictionary.	Char		
AE3_GRADE	Grade of Third Adverse Event	What was the grade of the third adverse event?	Num ###		
AE3_SYSTEMORGAN	System / Organ Class Affected by Third Adverse Event	Which system / organ classes were affected by the third adverse event? List each system / organ class, separated by a comma, using the list provided with this dictionary.	Char		
AE3_OUTCOME	Outcome of Third Adverse Event	What was the outcome of the third adverse event?	Category	Recov NoRecov Died	Recovered Not Recovered Died
7.L0_00100ML	Adverse Event	Advoise Event		Dieu	Dieu

Follow-Up Culture Results*								
Field	Variable	Additional Information	Format	Category Coding	Category Labeling			
		Market de la		Pos	Positive			
CULTURE MONTH	Culture Result	What is the culture result for the	Cotogoni	Neg	Negative			
CULTURE_MONTH1	Month 1	sputum sample tested during month 1?	Category	Contam	Contaminated			
		11		ND	Not Done			
		Mark to the endline many the feether		Pos	Positive			
OUT TUDE MONTUS	Culture Result	What is the culture result for the	0-1	Neg	Negative			
CULTURE_MONTH2	Month 2	sputum sample tested during month	Category	Contam	Contaminated			
		2?		ND	Not Done			
		140		Pos	Positive			
OUT TURE MONTUS	Culture Result	What is the culture result for the		Neg	Negative			
CULTURE_MONTH3	Month 3	sputum sample tested during month	Category	Contam	Contaminated			
		3?		ND	Not Done			
				Pos	Positive			
	Culture Result	What is the culture result for the		Neg	Negative			
CULTURE_MONTH4	Month 4	sputum sample tested during month	Category	Contam	Contaminated			
	Wionui 4	4?		ND	Not Done			
				Pos	Positive			
	Cultura Daguit	What is the culture result for the		Neg	Negative			
CULTURE_MONTH5	Culture Result	sputum sample tested during month	Category	Contam				
	Month 5	5?			Contaminated			
				ND	Not Done			
		What is the culture result for the		Pos	Positive			
CULTURE_MONTH6	Culture Result	sputum sample tested during month	Category	Neg	Negative			
OOLIGIKE_MONTIO	Month 6	6?	Category	Contam	Contaminated			
i e		0:		ND	Not Done			
		Addition the continue and the addition		Pos	Positive			
CHITHDE MONTHS	Culture Result	What is the culture result for the sputum sample tested during month 7?	0-4	Neg	Negative			
CULTURE_MONTH7	Month 7		Category	Contam	Contaminated			
				ND	Not Done			
		What is the culture result for the sputum sample tested during month		Pos	Positive			
	Culture Result			Neg	Negative			
CULTURE_MONTH8	Month 8		Category	Contam	Contaminated			
	Worter	8?		ND	Not Done			
		+		Pos	Positive			
	Culture Result Month 9	What is the culture result for the sputum sample tested during month 9?	Category	Neg	Negative			
CULTURE_MONTH9								
				Contam	Contaminated			
				ND	Not Done			
		What is the culture result for the sputum sample tested during month 10?	Category	Pos	Positive			
CULTURE_MONTH10	Culture Result			Neg	Negative			
	Month 10			Contam	Contaminated			
				ND	Not Done			
		What is the culture result for the		Pos	Positive			
CULTURE MONTH11	Culture Result	sputum sample tested during month	Category	Neg	Negative			
COLTORE_MONTHIT	Month 11	11?	Calegory	Contam	Contaminated			
		1119		ND	Not Done			
				Pos	Positive			
	Culture Result	What is the culture result for the			Negative			
CULTURE_MONTH12	Culture Result	sputum sample tested during month	Category	Neg	Negative Contaminated			
CULTURE_MONTH12	Culture Result Month 12		Category	Neg Contam	Contaminated			
CULTURE_MONTH12		sputum sample tested during month	Category	Neg Contam ND	Contaminated Not Done			
CULTURE_MONTH12	Month 12	sputum sample tested during month		Neg Contam ND Pos	Contaminated Not Done Positive			
CULTURE_MONTH12 CULTURE_MONTH13	Month 12 Culture Result	sputum sample tested during month 12?		Neg Contam ND Pos Neg	Contaminated Not Done Positive Negative			
	Month 12	sputum sample tested during month 12? What is the culture result for the	Category	Neg Contam ND Pos Neg Contam	Contaminated Not Done Positive Negative Contaminated			
	Month 12 Culture Result	sputum sample tested during month 12? What is the culture result for the sputum sample tested during month		Neg Contam ND Pos Neg Contam ND	Contaminated Not Done Positive Negative Contaminated Not Done			
	Month 12 Culture Result Month 13	sputum sample tested during month 12? What is the culture result for the sputum sample tested during month		Neg Contam ND Pos Neg Contam ND Pos	Contaminated Not Done Positive Negative Contaminated Not Done Positive			
	Month 12 Culture Result Month 13 Culture Result	sputum sample tested during month 12? What is the culture result for the sputum sample tested during month 13? What is the culture result for the	Category	Neg Contam ND Pos Neg Contam ND Pos Nheg ND Pos Neg ND Pos Neg	Contaminated Not Done Positive Negative Contaminated Not Done Positive Negative			
CULTURE_MONTH13	Month 12 Culture Result Month 13	sputum sample tested during month 12? What is the culture result for the sputum sample tested during month 13? What is the culture result for the sputum sample tested during month		Neg Contam ND Pos Neg Contam ND Pos Neg Contam ND Pos Neg Contam	Contaminated Not Done Positive Negative Contaminated Not Done Positive Negative Contaminated			
CULTURE_MONTH13	Month 12 Culture Result Month 13 Culture Result	sputum sample tested during month 12? What is the culture result for the sputum sample tested during month 13? What is the culture result for the	Category	Neg Contam ND Pos Neg Contam ND Pos Neg Contam ND Pos Neg Contam ND Neg	Contaminated Not Done Positive Negative Contaminated Not Done Positive Negative Contaminated Not Done Negative Contaminated Not Done			
CULTURE_MONTH13	Month 12 Culture Result Month 13 Culture Result	sputum sample tested during month 12? What is the culture result for the sputum sample tested during month 13? What is the culture result for the sputum sample tested during month 14?	Category	Neg Contam ND Pos Neg Contam ND Pos Neg Contam ND Pos Neg Contam	Contaminated Not Done Positive Negative Contaminated Not Done Positive Negative Contaminated			
CULTURE_MONTH13 CULTURE_MONTH14	Month 12 Culture Result Month 13 Culture Result	sputum sample tested during month 12? What is the culture result for the sputum sample tested during month 13? What is the culture result for the sputum sample tested during month 14? What is the culture result for the	Category	Neg Contam ND Pos Neg Contam ND Pos Neg Contam ND Pos Neg Contam ND Neg	Contaminated Not Done Positive Negative Contaminated Not Done Positive Negative Contaminated Not Done Negative Contaminated Not Done			
CULTURE_MONTH13	Month 12 Culture Result Month 13 Culture Result Month 14	sputum sample tested during month 12? What is the culture result for the sputum sample tested during month 13? What is the culture result for the sputum sample tested during month 14? What is the culture result for the sputum sample tested during month	Category	Neg Contam ND Pos Neg Contam ND Pos Neg Contam ND Pos Neg Contam ND Pos	Contaminated Not Done Positive Negative Contaminated Not Done Positive Negative Contaminated Not Done Positive Negative Contaminated Not Done Positive			
CULTURE_MONTH13 CULTURE_MONTH14	Month 12 Culture Result Month 13 Culture Result Month 14 Culture Result	sputum sample tested during month 12? What is the culture result for the sputum sample tested during month 13? What is the culture result for the sputum sample tested during month 14? What is the culture result for the	Category	Neg Contam ND Pos	Contaminated Not Done Positive Negative			
CULTURE_MONTH13 CULTURE_MONTH14	Month 12 Culture Result Month 13 Culture Result Month 14 Culture Result Month 15	sputum sample tested during month 12? What is the culture result for the sputum sample tested during month 13? What is the culture result for the sputum sample tested during month 14? What is the culture result for the sputum sample tested during month 15?	Category	Neg Contam ND Pos Neg Contam ND Pos Neg Contam ND Pos Neg Contam ND Pos Neg Contam ND ND Neg Contam ND ND Neg Contam ND	Contaminated Not Done Positive Negative Contaminated Not Done			
CULTURE_MONTH13 CULTURE_MONTH14	Month 12 Culture Result Month 13 Culture Result Month 14 Culture Result	sputum sample tested during month 12? What is the culture result for the sputum sample tested during month 13? What is the culture result for the sputum sample tested during month 14? What is the culture result for the sputum sample tested during month	Category	Neg Contam ND Pos Neg Contam ND Pos Neg Contam ND Pos Neg Contam ND Pos Neg Contam ND Contam ND Contam ND Contam ND Contam ND Contam	Contaminated Not Done Positive Negative Contaminated Contaminated			

CULTURE_MONTH17 Culture Result Month 17 Culture Result Month 17 Culture Result Month 18 Culture Result Month 18 Culture Result Month 18 Culture Result Month 19 Culture Result Month 20 Culture Result Month 20 Culture Result Month 20 Culture Result Month 20 Culture Result Month 21 What is the culture result for the sputum sample tested during month 21? Culture Result Month 21 Culture Result Month 22 Culture Result Month 23 What is the culture result for the sputum sample tested during month 21? Culture Result Month 23 What is the culture result for the sputum sample tested during month 21? Culture Result Month 23 Culture Result Month 23 What is the culture result for the sputum sample tested during month 21? Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 Culture Resul	Follow-Up Culture Results*							
CULTURE_MONTH17 Culture Result Month 17 Culture Result Month 17 Culture Result Month 18 Culture Result Month 19 Culture Result Month 20 Culture Result Month 20 Culture Result Month 20 Culture Result Month 21 Culture Result Month 22 Culture Result Month 23 What is the culture result for the sputum sample tested during month 21? What is the culture result for the sputum sample tested during month 22? What is the culture result for the sputum sample tested during month 22? What is the culture result for the sputum sample tested during month 22? What is the culture result for the sputum sample tested during month 22? Culture Result Month 23 What is the culture result for the sputum sample tested during month 22? Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 What is the culture result for the sputum sample tested during month 24? Culture Result Month 24 Culture Resul	Field	Variable	Additional Information	Format				
CULTURE_MONTH17 Culture Result Month 17 Culture Result Month 17 Culture Result Month 18 Culture Result Month 19 Culture Result Month 20 Culture Result Month 20 Culture Result Month 20 Culture Result Month 21 Culture Result Month 22 Culture Result Month 22 Culture Result Month 23 Culture Result Month 23 Culture Result Month 23 What is the culture result for the sputum sample tested during month 21? What is the culture result for the sputum sample tested during month 21? Culture Result Month 21 Culture Result Month 22 Culture Result Month 23 What is the culture result for the sputum sample tested during month 21? Culture Result Month 23 Culture Result Month 23 What is the culture result for the sputum sample tested during month 21? Culture Result Month 23 Culture Result Month 23 Culture Result Month 23 Culture Result Month 23 Culture Result Month 24 Culture					ND	Not Done		
CULTURE_MONTH17 CULTURE_MONTH18 CULTURE_MONTH18 CULTURE_MONTH18 CULTURE_MONTH19 CULTURE_MONTH19 CULTURE_MONTH20 CULTURE_MONTH21 CULTURE_MONTH21 CULTURE_MONTH21 CULTURE_MONTH21 CULTURE_MONTH21 CULTURE_MONTH22 CULTURE_MONTH22 CULTURE_MONTH22 CULTURE_MONTH24 CULTURE_MONTH25 CULTURE_MONTH25 CULTURE_MONTH26 CULTURE_MONTH26 CULTURE_MONTH27 CULTURE_MONTH27 CULTURE_MONTH28 CULTURE_MONTH29 CULTUR					Pos	Positive		
CULTURE_MONTH18 Culture Result Month 18 Culture Result Month 19 Culture Result Month 20 Culture Result Month 21 Culture Result Month 22 Culture Result Month 22 Culture Result Month 22 Culture Result Month 22 Culture Result Month 23 Culture Result Month 23 Culture Result Month 23 What is the culture result for the sputum sample tested during month 22? Culture Result Month 23 What is the culture result for the sputum sample tested during month 22? Culture Result Month 23 What is the culture result for the sputum sample tested during month 22? Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 Culture Result Month 26 Culture Result Month 27 Culture Result Month 28 Culture Result Month 29 Culture Res	CHITHDE MONTHAZ	Culture Result		Cotogoni	Neg	Negative		
CULTURE_MONTH18 Culture Result Month 18 Culture Result Month 18 Culture Result Month 19 Culture Result Month 19 Culture Result Month 19 Culture Result Month 20 Culture Result Month 20 Culture Result Month 20 Culture Result Month 21 Culture Result Month 22 Culture Result Month 22 Culture Result Month 23 What is the culture result for the sputum sample tested during month 22? What is the culture result for the sputum sample tested during month 22? Culture Result Month 23 What is the culture result for the sputum sample tested during month 22? What is the culture result for the sputum sample tested during month 23? What is the culture result for the sputum sample tested during month 23? Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 What is the culture result for the sputum sample tested during month 24? Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg	COLTURE_MONTHI7	Month 17		Category	Contam	Contaminated		
CULTURE_MONTH18 Culture Result Month 18 Culture Result Month 18 Culture Result Month 18 Culture Result Month 19 Culture Result Month 20 Culture Result Month 21 Culture Result Month 22 Culture Result Month 22 Culture Result Month 22 Culture Result Month 22 Culture Result Month 23 What is the culture result for the sputum sample tested during month 22? What is the culture result for the sputum sample tested during month 22? Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? Culture Result Month 23 Culture Result Month 24 What is the culture result for the sputum sample tested during month 23? Culture Result Month 23 Culture Result Month 24 Culture Result Month 24 Culture Result Month 24 Culture Result Month 26 Culture Result Month 27 Category Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminat			17 !		ND	Not Done		
CULTURE_MONTH18 Culture Result Month 18 Culture Result Month 18 Culture Result Month 18 Culture Result Month 19 Culture Result Month 20 Culture Result Month 21 Culture Result Month 22 Culture Result Month 23 Culture Result Month 23 Culture Result Month 24 Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 What is the culture result for the sputum sample tested during month 24 What is the culture result for the sputum sample tested during month 24 Culture Result			M/h at is the anothern manual for the		Pos	Positive		
CULTURE_MONTH19 Culture Result Month 19 Culture Result Month 19 Culture Result Month 19 Culture Result Month 19 Culture Result Month 20 Culture Result Month 21 Culture Result Month 22 Culture Result Month 23 Culture Result Month 24 Culture Result Month 29 Culture Result Month 20 Culture Result Month 20 Culture Result Month 20 Culture Result Month 20 Cult	CHITHE MONTHA	Culture Result		Cotogoni	Neg	Negative		
CULTURE_MONTH19 Culture Result Month 19 Culture Result Month 20 Culture Result Month 21 Culture Result Month 22 Culture Result Month 23 Culture Result Month 23 Culture Result Month 24 Culture Result Month 23 Culture Result Month 23 Culture Result Month 23 Culture Result Month 24 Culture Result Month 26 Culture Result Month 27 Culture Result Month 28 Culture Result Month 29 Cult	CULTURE_MONTH16	Month 18	, , ,	Category	Contam	Contaminated		
CULTURE_MONTH19 Culture Result Month 19 Culture Result Month 19 Culture Result Month 20 Culture Result Month 21 Culture Result Month 22 Culture Result Month 23 Culture Result Month 24 Culture Result Month 26 Culture Result Month 27 Culture Result Month 28 Culture Result Month 29 Cult			10?		ND	Not Done		
CULTURE_MONTH20 Culture Result Month 19 Culture Result Month 20 Culture Result Month 20 Culture Result Month 20 Culture Result Month 20 Culture Result Month 21 Culture Result Month 22 Culture Result Month 23 Culture Result Month 24 Culture Result Month 26 Culture Result Month 27 Culture Result Month 28 Culture Result Month 29 Cult			NA/least in the analysis was used for the		Pos	Positive		
CULTURE_MONTH20 Culture Result Month 20 Culture Result Month 20 Culture Result Month 20 Culture Result Month 20 Culture Result Month 21 Culture Result Month 22 Culture Result Month 23 Culture Result Month 23 What is the culture result for the sputum sample tested during month 22? What is the culture result for the sputum sample tested during month 22? Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? What is the culture result for the sputum sample tested during month 23? What is the culture result for the sputum sample tested during month 23? What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 What is the culture result for the sputum sample tested during month 23? Category Category Category Category Category Category Category Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated Contaminated Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated Co	CHITHE MONTHA	Culture Result		Cotogoni	Neg	Negative		
CULTURE_MONTH20 Culture Result Month 20 Culture Result Month 21 Culture Result Month 22 Culture Result Month 23 What is the culture result for the sputum sample tested during month 22? What is the culture result for the sputum sample tested during month 22? Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? Culture Result Month 23 Culture Result Month 24 What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 Culture Result Month 26 Culture Result Month 27 Category Material Month Mon	COLTURE_MONTH19	Month 19		Category	Contam	Contaminated		
CULTURE_MONTH20 Culture Result Month 20 Culture Result Month 20 Culture Result Month 20 Culture Result Month 21 Culture Result Month 22 Culture Result Month 23 Culture Result Month 24 Culture Result Month 26 Culture Result Month 27 Culture Result Month 28 Culture Result Month 29 Cult					ND	Not Done		
CULTURE_MONTH20 Culture Result Month 20 Culture Result Month 21 Culture Result Month 22 Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 Culture Result Month 24 What is the culture result for the sputum sample tested during month 23? What is the culture result for the sputum sample tested during month 24? Culture Result Month 24 Culture Result Month 26 Culture Result Month 27 Category Month			sputum sample tested during month		Pos	Positive		
CULTURE_MONTH21 Culture Result Month 21 Culture Result Month 22 What is the culture result for the sputum sample tested during month 22? Culture Result Month 23 Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated Cont	OLU TURE MONTURO			Cotogoni	Neg	Negative		
CULTURE_MONTH21 Culture Result Month 21 Culture Result Month 21 Culture Result Month 21 Culture Result Month 21 Culture Result Month 22 Culture Result Month 22 Culture Result Month 22 Culture Result Month 22 Culture Result Month 23 Culture Result Month 24 Culture Result Month 25 Culture Result Month 26 Culture Result Month 27 Category Month	CULTURE_MONTH20			Category	Contam	Contaminated		
CULTURE_MONTH21 Culture Result Month 21 Culture Result Month 21 Culture Result Month 21 Culture Result Month 21 Culture Result Month 22 Culture Result Month 23 Culture Result Month 24 Culture Result Month 25 Culture Result Month 26 Culture Result Month 27 Category Meg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive ND Not Done ND Not Done Culture Result ND Not Done Pos Positive ND Not Done ND Not Done Culture Result ND Not Done ND Not Done ND Not Done Culture Result ND Not Done					ND	Not Done		
CULTURE_MONTH21 Culture Result Month 21 Sputum sample tested during month 21? Culture Result Month 21 Culture Result Month 22 Culture Result Month 22 Culture Result Month 22 What is the culture result for the sputum sample tested during month 22? Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 Category Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Not Done Culture Result ND Not Done Pos Positive Not Done Contam Contaminated ND Not Done Pos Positive Not Done Contam Contaminated ND Not Done Pos Positive Contam Contaminated ND Not Done Contam Contaminated ND Not Done Pos Positive Not Positive Not			sputum sample tested during month	Catagoni	Pos	Positive		
CULTURE_MONTH22 Culture Result Month 22 Culture Result Month 22 Culture Result Month 22 Culture Result Month 22 Culture Result Month 23 Culture Result Month 24 Culture Result Month 25 Culture Result Month 26 Culture Result Month 27 Category Month	CULTURE MONTHS	Culture Result			Neg	Negative		
CULTURE_MONTH22 Culture Result Month 22 Culture Result Month 22 What is the culture result for the sputum sample tested during month 22? What is the culture result for the sputum sample tested during month 22? Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? Culture Result Month 23 Culture Result Month 24 What is the culture result for the sputum sample tested during month 24 Culture Result Month 25 Category Contam Contaminated ND Not Done Pos Positive ND Not Done Pos Positive ND Not Done Category Contam Contaminated ND Not Done Pos Positive ND Not Done Contam Contaminated	CULTURE_MONTH21	Month 21		Category	Contam	Contaminated		
CULTURE_MONTH22 Culture Result Month 22 Culture Result Month 22 Culture Result Month 22 Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? Culture Result Month 23 Culture Result Month 24 What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 Category Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Culture Result Not Done Pos Positive Not Done Culture Result Not Done Pos Positive Neg Negative Contam Contaminated Not Done Pos Positive Not Done Culture Result Not Done Pos Positive Neg Negative Contam Contaminated Not Done Pos Positive Not Done Culture Result Not Done Pos Positive Neg Negative Contam Contaminated Not Done Pos Positive Not Done Pos P					ND	Not Done		
CULTURE_MONTH22 Culture Result Month 22 Sputum sample tested during month 22? Culture Result Month 23 Culture Result Month 24 Culture Result Month 25 Culture Result Month 26 Culture Result Month 27 Category Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Pos Positive Neg Negative Contam Contaminated ND Not Done Culture Result Month 24 Culture Result Month 25 Culture Result Month 26 Culture Result Month 27 Category Month Mon			M/h at is the anothern manual for the		Pos	Positive		
CULTURE_MONTH23 Culture Result Month 23 Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? Culture Result Month 24 Culture Result Month 23 Category Neg Negative Month ND Not Done Pos Positive ND Not Done Contam Contaminated Contam Contaminated Contam Contaminated	CHITHE MONTHS	Culture Result		Cotogoni	Neg	Negative		
CULTURE_MONTH23 Culture Result Month 23 Culture Result Month 23 Culture Result Month 23 Culture Result Month 23 Culture Result Month 24 Culture Result Month 25 Culture Result Month 26 Culture Result Month 27 Culture Result Month 28 Culture Result Month 29 Cult	CULTURE_MONTH22	Month 22		Category	Contam	Contaminated		
CULTURE_MONTH23 Culture Result Month 23 What is the culture result for the sputum sample tested during month 23? Culture Result Month 23 Culture Result Month 24 Culture Result Month 24 What is the culture result for the sputum sample tested during month 24? What is the culture result for the sputum sample tested during month 24? Category Contam Contaminated ND Neg Negative Neg Neg Negative Contam Contaminated			22!		ND	Not Done		
CULTURE_MONTH23 Culture Result Month 23 Sputum sample tested during month 23? Sputum sample tested during month 23? Category Contam Contaminated ND Not Done What is the culture result for the sputum sample tested during month 24 Culture Result Month 24 Culture Result Month 24 Culture Result Month 24 Culture Result Sputum sample tested during month 24 Category Category Neg Negative Not Done Pos Positive Neg Negative Contam Contaminated			M/h at is the anothern manual for the		Pos	Positive		
23? 23? Culture Result Month 24 Culture Result Tonth Sputum sample tested during month Category Contam Contaminated ND Not Done What is the culture result for the sputum sample tested during month Category Contam Contaminated ND Not Done Culture Result Month 24 Category Contam Contaminated ND Not Done	CULTURE MONTHS	Culture Result		0-4	Neg	Negative		
CULTURE_MONTH24 Culture Result Month 24 Culture Result Month 24 Culture Result Month 24 Culture Result Month 24 Culture Result Sputum sample tested during month 24 Culture Result Sputum sample tested during month 24 Category Contam Contaminated	CULTURE_MONTH23	Month 23	, , ,	Category	Contam	Contaminated		
CULTURE_MONTH24 Culture Result Month 24 Culture Result Month 24 Culture Result Sputum sample tested during month 24 Category Contam Contaminated			23?		ND	Not Done		
CULTURE_MONTH24 Culture Result Month 24 sputum sample tested during month Category Contam Contaminated Neg Negative Contam Contaminated Contam Contaminated Contam Contaminated Contam Contaminated Contaminated			Milest is the soultime result for the		Pos			
Month 24 Sputum sample tested during month Category Contam Contaminated	CHITHDE MONTHS4	Culture Result		0-1	Neg	Negative		
Z4 ! ND Not Dono	CULTURE_MONTH24	Month 24		Category	Contam	Contaminated		
			24!		ND	Not Done		

^{*}Month 1 refers to the result of the sample taken between day 31 and 60 that is closest to day 31 and valid (i.e., Positive or Negative); Month 2 refers to the sample taken between day 61 and 90 that is closest to day 61 and valid (i.e., Positive or Negative); the remaining months follow the same pattern. Any MTB colonies seen should be considered positive. If multiple samples are taken on a given day, a positive-dominant approach should be taken, whereby a patient is positive if a single positive sample is found. A patient sample should only be classified as contaminated if all samples from that month were contaminated.

Follow-Up Smear Microscopy Results*								
Field	Variable	Additional Information	Format	Category Coding	Category Labeling			
		What is the smear result for the		Pos	Positive			
SMEAR_MONTH1	Smear Result	sputum sample tested during month	Category	Neg	Negative			
OWE, INC. INC. ITT	Month 1	1?	Catogory	Contam	Contaminated			
				ND	Not Done			
		What is the smear result for the		Pos	Positive			
SMEAR_MONTH2	Smear Result	sputum sample tested during month	Category	Neg	Negative			
_	Month 2	2?		Contam ND	Contaminated Not Done			
_				Pos	Positive			
	Smear Result	What is the smear result for the		Neg	Negative			
SMEAR_MONTH3	Month 3	sputum sample tested during month	Category	Contam	Contaminated			
	Widititi	3?		ND	Not Done			
				Pos	Positive			
	Smear Result	What is the smear result for the		Neg	Negative			
SMEAR_MONTH4	Month 4	sputum sample tested during month	Category	Contam	Contaminated			
	Wionan i	4?		ND	Not Done			
				Pos	Positive			
0.45.6	Smear Result	What is the smear result for the		Neg	Negative			
SMEAR_MONTH5	Month 5	sputum sample tested during month	Category	Contam	Contaminated			
		5?		ND	Not Done			
				Pos	Positive			
ONE AD MONETUS	Smear Result	What is the smear result for the		Neg	Negative			
SMEAR_MONTH6	Month 6	sputum sample tested during month	Category	Contam	Contaminated			
		6?		ND	Not Done			
	Smear Result	140 41 41		Pos	Positive			
CNAFAD MONITUR		What is the smear result for the sputum sample tested during month 7?	Category	Neg	Negative			
SMEAR_MONTH7	Month 7			Contam	Contaminated			
				ND	Not Done			
		What is the smear result for the sputum sample tested during month 8?	Category	Pos	Positive			
CATE A D. MONTHIO	Smear Result			Neg	Negative			
SMEAR_MONTH8	Month 8		Category	Contam	Contaminated			
				ND	Not Done			
	Smear Result	What is the smear result for the sputum sample tested during month 9?		Pos	Positive			
SMEAR_MONTH9			Category	Neg	Negative			
SWEAK_WONTTIS	Month 9			Contam	Contaminated			
				ND	Not Done			
		What is the smear result for the	Category	Pos	Positive			
SMEAR MONTH10	Smear Result	sputum sample tested during month		Neg	Negative			
OME/ IT _ INGITITIO	Month 10			Contam	Contaminated			
				ND	Not Done			
		What is the smear result for the		Pos	Positive			
SMEAR MONTH11	Smear Result	sputum sample tested during month	Category	Neg	Negative			
	Month 11	11?		Contam	Contaminated			
				ND	Not Done			
	0 5 1	What is the smear result for the		Pos	Positive			
SMEAR_MONTH12	Smear Result	sputum sample tested during month	Category	Neg	Negative			
	Month 12	12?		Contam ND	Contaminated			
_				Pos	Not Done Positive			
	Smear Result	What is the smear result for the		Neg	Negative			
SMEAR_MONTH13	Month 13	sputum sample tested during month	Category	Contam	Contaminated			
	IVIOITIII 13	13?		ND	Not Done			
				Pos	Positive			
	Smear Result	What is the smear result for the		Neg	Negative			
SMEAR_MONTH14	Month 14	sputum sample tested during month	Category	Contam	Contaminated			
	Mondi IT	14?		ND	Not Done			
	+			Pos	Positive			
	Smear Result	What is the smear result for the		Neg	Negative			
SMEAR_MONTH15	Month 15	sputum sample tested during month	Category	Contam	Contaminated			
	Wionan 10	15?		ND	Not Done			
	Smear Result		1_	Pos	Positive			
SMEAR_MONTH16	Month 16		Category	Neg	Negative			
		1	1	1				

Follow-Up Smear Microscopy Results*							
Field	Variable	Additional Information	Format	Category Coding	Category Labeling		
		What is the smear result for the		Contam	Contaminated		
		sputum sample tested during month 16?		ND	Not Done		
		Mile at in the angree of manufacture for the		Pos	Positive		
CMEAD MONTHAT	Smear Result	What is the smear result for the	Cotogomi	Neg	Negative		
SMEAR_MONTH17	Month 17	sputum sample tested during month 17?	Category	Contam	Contaminated		
		17 !		ND	Not Done		
		Mile at in the angree of manufacture for the		Pos	Positive		
CMEAD MONTHA	Smear Result	What is the smear result for the	Cotogomi	Neg	Negative		
SMEAR_MONTH18	Month 18	sputum sample tested during month 18?	Category	Contam	Contaminated		
		10?		ND	Not Done		
		Miles Carlos and a second for the		Pos	Positive		
CMEAD MONTHA	Smear Result Month 19	What is the smear result for the sputum sample tested during month 19?	Category	Neg	Negative		
SMEAR_MONTH19				Contam	Contaminated		
				ND	Not Done		
		What is the smear result for the sputum sample tested during month 20?		Pos	Positive		
CMEAD MONTHS	Smear Result Month 20		Cotogomi	Neg	Negative		
SMEAR_MONTH20			Category	Contam	Contaminated		
				ND	Not Done		
		140	Category	Pos	Positive		
CMEAD MONTHS	Smear Result	What is the smear result for the		Neg	Negative		
SMEAR_MONTH21	Month 21	sputum sample tested during month 21?		Contam	Contaminated		
				ND	Not Done		
		Mile at in the angree of manufacture for the		Pos	Positive		
CMEAD MONTHS	Smear Result	What is the smear result for the	Cotogomi	Neg	Negative		
SMEAR_MONTH22	Month 22	sputum sample tested during month 22?	Category	Contam	Contaminated		
		22!		ND	Not Done		
		Miles the second second for the		Pos	Positive		
CMEAD MONTHS	Smear Result	What is the smear result for the	Cotogomi	Neg	Negative		
SMEAR_MONTH23	Month 23	sputum sample tested during month 23?	Category	Contam	Contaminated		
		23!		ND	Not Done		
		Mhat is the amount regult for the		Pos	Positive		
CMEAD MONTHS	Smear Result	What is the smear result for the	Category	Neg	Negative		
SMEAR_MONTH24	Month 24	sputum sample tested during month 24?		Contam	Contaminated		
		Z4:		ND	Not Done		

^{*}Any acid-fast bacilli seen should be considered positive. Month 1 refers to the result of the sample taken between day 31 and 60 that is closest to day 31 and valid (i.e., Positive or Negative); Month 2 refers to the sample taken between day 61 and 90 that is closest to day 61 and valid (i.e., Positive or Negative); the remaining months follow the same pattern. If multiple samples are taken within a given day, a positive-dominant approach should be taken, whereby a patient is positive if a single positive sample is found. A patient sample should be classified as contaminated only if all samples from that month were contaminated.

	Treatment Outcome Information							
Field	Variable	Additional Information	Format	Category Coding	Category Labeling			
OUTCOME_DEFINITION	End-of-Treatment Outcome	Specify the guideline year the outcome definition follows—this is preferably	Category	WHO2013	2013 Definitions			
COTCOME_BET INTHON	Definition	the 2013 guidelines but can follow 2005 guidelines if not available.	Calegory	WHO2005	2005 Definitions			
				Cure	Cure			
		End of treatment outcome		Complete	Treatment Complete			
OUTCOME	End-of-Treatment Outcome	assigned to the patient, following the outcome year	Category	Fail	Treatment Failure			
		specified above.		Death	Death			
				LTFU	Loss to Follow- Up			
		Did the patient culture convert (defined as two		Υ	Yes			
CULTURECONV*	Culture	consecutive negative cultures taken at least 28 d	Cotogoni	N	No			
COLTURECONV	Conversion	apart)? If the patient was culture negative at baseline, list as BaseNeg.	Category	BaseNeg	Baseline Negative			
CULTURECONV_DATE	Date of Culture Conversion	If the patient culture converted, what was the date of conversion (defined as the date of the first of the two consecutive negative cultures)?	Date					
	Culture Conversion by	If exact date of conversion is		Υ	Yes			
TWOCONV		unknown, did culture conversion occur before the	Category	N	No			
	Month Two	end of month two?		U	Unknown			
	Culture	If exact date of conversion is unknown, did culture		Y N	Yes No			
SIXCONV	Conversion by Month Six	conversion occur before the end of month six?	Category	U	Unknown			
		If patient converted or was		Υ	Yes			
		culture negative at baseline, was there culture reversion		N	No			
CULTUREREV*	Culture Reversion	(defined as two consecutive positive cultures taken at	Category	U	Unknown			
CULTUREREV_DATE	Date of Culture Reversion	least 28 d apart)? If patient had culture reversion, what was the date of reversion (defined as the date of the first of the two consecutive positive cultures)?	Date					
RECURRENCE_MONITORING	Post-Treatment Recurrence Monitoring	Was post-treatment monitoring for recurrence performed?	Category	Y N	Yes No			
RECURRENCE_FOLLOWUP_DUR	Duration of Recurrence Monitoring	What was the duration of recurrence monitoring, in months?	Num ###					
RECURRENCE_OUTCOME	Occurrence of	Did the patient experience recurrence?	Category	Y	Yes			
RECURRENCE_DATE	Recurrence Date of Recurrence	What was the date of the recurrence episode?	Date	114	No			
	recuirence	If resources permitted, was		Relapse	Relapse			
RELAPSE_REINFECT	Relapse or	the recurrence classified as	Category	Reinfect	Reinfection			
NELAI OL_ILLINI LOI	Reinfection	a true relapse or as a	Calegory	U	Unknown			
*These can be reported by the individual pr	oviding data or calculate	reinfection?	multiple culti	ros takon at th				

^{*}These can be reported by the individual providing data or calculated by an analyst. In the instance of multiple cultures taken at the same time, a positive dominant approach should be taken, i.e., the result should be considered positive if any of the samples are positive. In the case of contaminated results, these should be discarded when calculating time to culture conversion or reversion.

Drug Abbreviations, System/Organ Classes, and End-of-Treatment Outcome Definitions

The tables contained within this section are intended to promote standardization in coding of drugs, outcomes, and adverse events.

Tanada and and and and and and and and an	breviation
Isoniazid H	
Rifampin R	
Ethambutol E	
Pyrazinamide Z	
High Dose Isoniazid Hig	hH
Streptomycin S	
Rifabutin Rfb	
Amikacin Am	<u> </u>
Capreomycin Cm	
Kanamycin Km	l .
Ofloxacin Ofx	
Ciprofloxacin	
Moxifloxacin Mfx	(
Levofloxacin Lfx	
Gatifloxacin Gfx	(
Sparfloxacin Sfx	
Ethionamide Eto)
Prothionamide Pto)
Cycloserine Cs	
Terizidone Trd	
Para-Aminosalicylic Acid PAS	S
Linezolid Lzd	1
Clofazimine	
Amoxicillin and Clavulanic Acid Am	xClv
Imipenem-Cilastatin Ipm	1
Meropenem Mpi	m
Bedaquiline Bdd	9
Delamanid Dlm	n
Pretomanid Pa	
Perchlozone Pcz	7
Thioacetazone T	
Rifapentine Rpt	t
Second Line Injectables SLI	
Fluoroquinolones FQ	

Drug Name / Drug Class of Antiretroviral Therapy	Abbreviation
Nucleoside/Nucleotide Reverse transcription Inhibitor	NRTI
Abacavir	ABC
Didanosine	ddl
Emtricitabine	FTC
Lamivudine	3TC
Stavudine	d4T
Tenofovir alafenamide	TAF
Tenofovir disoproxil fumarate	TDF
Zidovudine	AZT or ZDV
Non-nucleoside Reverse transcription Inhibitor	NNRTI
Delaviridine	DLV
Efavirenz	EFV
Etavirine	ETR
Nevirapine	NVP
Rilpivirine	RPV
Protease Inhibitor	PI
Amprenavir	AMV
Atazanavir	ATV
Darunavir	DRV
Fosamprenavir	FPV
Indinavir	IDV
Lopinavir + ritonavir	LPV/r
Nelfinavir	NFV
Saquinavir	SQV
Tipranavir	TPV
Fusion Inhibitor	FI
Enfuviritide	ENF or T-20
CCR5 Antagonist	CCR5
Maraviroc	MVC
Integrase Inhibitor	II
Bictegravir	BIC
Dolutegravir	DTG
Elvitegravir	EVG
Raltegravir	RAL

SYSTEM/ORGAN CLASS
Blood and lymphatic system disorders
Cardiac disorders
Congenital, familial and genetic disorders
Ear and labyrinth disorders
Endocrine disorders
Eye disorders
Gastrointestinal disorders
General disorders and administration site conditions
Hepatobiliary disorders
Immune system disorders
Infections and infestations
Injury, poisoning and procedural complications
Investigations
Metabolism and nutrition disorders
Musculoskeletal and connective tissue disorders
Neoplasms benign, malignant and unspecified (incl cysts and polyps)
Nervous system disorders
Pregnancy, puerperium and perinatal conditions
Psychiatric disorders
Renal and urinary disorders
Reproductive system and breast disorders
Respiratory, thoracic and mediastinal disorders
Skin and subcutaneous tissue disorders
Social circumstances
Surgical and medical procedures
Vascular disorders

WHO 2013 Outcome Definitions (Preferred)		
Outcome	Definition	
Cure	Treatment completed as recommended by the national policy without evidence of failure AND three or more consecutive cultures taken at least 30 d apart are negative after the intensive phase (or Month 8 if no intensive phase).	
Complete	Treatment completed as recommended by the national policy without evidence of failure BUT no record that three or more consecutive cultures taken at least 30 d apart are negative after the intensive phase (or Month 8 if no intensive phase).	
Failure*	Treatment terminated or need for permanent regimen change of at least two anti-TB drugs because of: (1) lack of conversion by the end of the intensive phase, or (2) bacteriological reversion in the continuation phase after conversion to negative, or (3) evidence of additional acquired resistance to fluoroquinolones or second-line injectable drugs, or (4) adverse drug reactions.	
Death	A patient who dies for any reason during the course of treatment	
Lost to Follow-up	A patient whose treatment was interrupted for 2 consecutive months or more.	

WHO 2005 (Laserson) Outcome Definitions (if 2013 not possible)		
Outcome	Definition	
Cure	Completed treatment according to program protocol and has at least five consecutive negative cultures from samples collected at least 30 d apart in the final 12 mo of treatment. If only one positive culture is reported during that time, and there is no concomitant clinical evidence of deterioration, a patient may still be considered cured, provided that this positive culture is followed by a minimum of three consecutive negative cultures taken at least 30 d apart.	
Complete	Completed treatment according to program protocol but does not meet the definition for cure because of lack of bacteriological results (i.e., fewer than five cultures were performed in the final 12 mo of treatment).	
Failure	Treatment will be considered to have failed if two or more of the five cultures recorded in the final 12 mo of therapy are positive, or if any one of the final three cultures is positive. (Treatment will also be considered to have failed if a clinical decision has been made to terminate treatment early because of poor clinical or radiological response or adverse events).	
Death	A patient who dies for any reason during the course of MDR/RR-TB treatment	
Lost to Follow-up	A patient whose treatment was interrupted for two or more consecutive months for any reason without medical approval.	

Example of an Initial Data Sharing Agreement (Can Be Modified on a Case-By-Case Basis)

LETTER OF AGREEMENT for IPD in MDR/RR TB

This letter of agreement is between the McGill University group (hereafter referred to as the McGill group) for an Individual Patient Data (IPD) meta-analysis in multidrug-resistant tuberculosis TB (MDR-TB), and [INSERT NAME OF INVESTIGATOR AND INSTITUTION] (hereafter referred to as the investigator), regarding the transfer and use of data collected by the investigator. The McGill group and the investigator agree to collaborate on [INSERT NAME OF PROJECT] according to the terms in this letter and those set out in the full project protocol, which is attached as Annex 1.

The McGill group agrees to:

- Obtain approval from the Research Ethics Board of the Montreal Chest Institute,
 McGill University Health Center for this research.
- Respect the confidentiality of all data received. They will not attempt to identify patients, nor contact patients directly.
- Respect the principle that the investigator continues to 'own' the data sent for inclusion in this analysis. When the data set is "cleaned" and preliminary analyses completed, a copy of the data set will be returned to the investigator.
- Perform data analysis that addresses the objectives specified in the attached study protocol only. Any additional analysis will be performed only after it has been approved by the investigator. For additional analyses that are closely related to these objectives, the investigator will be informed; approval will be assumed if the investigator does not reply within a specified interval. If the investigator has concerns or objections to any new analyses, these will be addressed and resolved before proceeding. Analyses to address completely novel objectives that have not been foreseen in the current study protocol must be actively approved by the investigator before these analyses are undertaken.
- Finish analyses and return the data to the investigator by the sunset date. This date will be the date by which the analyses must be completed, and any manuscript(s)

prepared. The tentative sunset date to complete analyses, and prepare related manuscripts is [INSERT DATE]. If a manuscript is submitted, the data must be held until peer review is completed, and then up to 1 year after publication – to allow time for responses to the findings (e.g., letters to the editor). However, after the sunset date no further new analysis can begin without agreement to the extension of the sunset date by the investigator.

- Share results of analyses with the investigator, and all members of the IPD group at intervals described in the study protocol.
- Prepare draft and final reports of results for the project and prepare manuscript(s) of results for publication. All draft reports and manuscripts will be reviewed and approved by the investigator, and all members of the IPD group before submission. The authorship of these reports will be "The Collaborative Group for Meta-Analysis of Updated Individual Patient Data in MDR-TB", followed by a listing of all members in alphabetic order. The corresponding author will be Dr. Menzies of McGill.

The investigator agrees to:

- Verify whether they require approval from their local Research Ethics Board, depending on their institution's policy. If so, the investigator will obtain this approval before sending the data to the McGill group. No additional data will be collected from the patients, thus investigator will not need to obtain patients' consent for this analysis.
- Transfer a data file of information on all patients who were members of a cohort of MDR-TB patients which the investigator reported in earlier publications. This patient dataset will be rendered completely anonymous before forwarding this to the Montreal Chest Institute by removing all personal identifiers.
- Become a member of The Collaborative Group for Meta-Analysis of Updated
 Individual Patient Data in MDR-TB. This Collaborative Group will review all
 preliminary and final results of analyses performed by the McGill group, as well

as all reports of results – for the guideline groups, for public presentation, and for publication.
Treat these preliminary results confidentially. The investigator will not publish

Treat these preliminary results confidentially. The investigator will not publish
 (including posting on the Internet), present in any public forum, nor disseminate
 through any media these results without approval from the McGill Group and
 other members of the IPD Collaborative Group.

Dr. Dick Menzies (for the McGill University Group)	Date
[Insert name and institution]	Date